

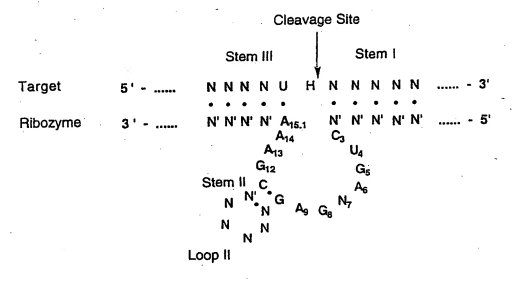
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(54) Title: COMPOSITIONS AND METHOD FOR MODULATION OF ALKALOID BIOSYNTHESIS AND FLOWER FORMATION IN PLANTS



(57) Abstract

An enzymatic nucleic acid molecule with RNA cleaving activity, wherein the nucleic acid molecule modulates the expression of a gene involved in the biosynthesis of alkaloid compounds and flower formation in a plant. A transgenic plant comprising nucleic acids encoding for an enzymatic nucleic acid molecule with RNA cleaving activity, wherein the nucleic acid molecule modulates the expression of a gene involved in the biosynthesis of alkaloid compounds of flower formation in a plant. An enzymatic nucleic acid molecule with RNA cleaving activity, wherein the nucleic acid molecule modulates the expression of solanidine UDP-glucose glucosyl-transferase gene or citrate synthase in plants.

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DESCRIPTION

Compositions And Method For Modulation Of Alkaloid Biosynthesis And Flower Formation In Plants

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Background of the Invention

The present invention concerns compositions and methods for the modulation of gene expression in plants, specifically using enzymatic nucleic acid molecules.

The following is a brief description of regulation of gene expression in plants. The discussion is not meant to be complete and is provided only for understanding of the invention that follows. This summary is not an admission that any of the work described below is prior art to the claimed invention.

There are a variety of strategies for modulating gene expression in plants. Traditionally, antisense RNA (reviewed in Bourque, 1995 Plant Sci 105, 125-149) and co-suppression (reviewed in Jorgensen, 1995 Science 268, 686-691) approaches have been used to modulate gene expression. Insertion mutagenesis of genes have also been used to silence gene expression. This approach, however, cannot be designed to specifically inactivate the gene of interest. Applicant believes that ribozyme technology offers an attractive new means to alter gene expression in plants.

Naturally occurring antisense RNA was first discovered in bacteria over a decade ago (Simons and Kleckner, 1983 Cell 34, 683-691). It is thought to be one way in which bacteria can regulate their gene expression (Green et al., 1986 Ann. Rev. Biochem. 55: 567-597; Simons 1988 Gene 72: 35-44). The first demonstration of antisense-mediated inhibition of gene expression was reported in mammalian cells (Izant and Weintraub

1984 <u>Cell</u> 36: 1007-1015). There are many examples in the literature for the use of antisense RNA to modulate gene expression in plants. Following are a few examples:

Shewmaker et al., U.S. Patent Nos. 5,107,065 and 5, 5 453,566 disclose methods for regulating gene expression in plants using antisense RNA.

It has been shown that an antisense gene expressed in plants can act as a dominant suppressor gene. Transgenic potato plants have been produced which express RNA antisense to potato or cassava granule bound starch In both of these cases, transgenic synthase (GBSS). plants have been constructed which have reduced or no GBSS activity or protein. These transgenic plants give rise to potatoes containing starch with dramatically 15 reduced amylose levels (Visser et al. 1991, Mol. Gen. Genet. 225: 2889-296; Salehuzzaman et al. 1993, Plant Mol. Biol. 23: 947-962).

Kull et al., 1995, J. Genet. & Breed. 49, 69-76 reported inhibition of amylose biosynthesis in tubers 20 from transgenic potato lines mediated by the expression of antisense sequences of the gene for granule-bound starch synthase (GBSS). The authors, however, indicated a failure to see any in vivo activity of ribozymes targeted against the GBSS RNA.

Antisense RNA constructs targeted against desaturase enzyme in canola have been shown to increase the level of stearic acid (C18:0) from 2% to 40% (Knutzon et. al., 1992 Proc. Natl. Acad. Sci. 89, 2624). was no decrease in total oil content or germination 30 efficiency in one of the high stearate lines. recent reviews are available which illustrate the utility of plants with modified oil composition (Ohlrogge, J. B. 1994 Plant Physiol. 104, 821; Kinney, A. J. 1994 Curr.

Opin. Cell Biol. 5, 144; Gibson et al. 1994 Plant Cell Envir. 17, 627).

inactivation was Homologous transgene documented in plants as an unexpected result of inserting 5 a transgene in the sense orientation and finding that both the gene and the transgene were down-regulated (Napoli et al., 1990 <u>Plant Cell</u> 2: 279-289). appears to be at least two mechanisms for inactivation of genetic sequences. One appears homologous 10 transcriptional inactivation via methylation, where duplicated DNA regions signal endogenous mechanisms for gene silencing. This approach of gene modulation involves either the introduction of multiple copies of transgenes or transformation of plants with transgenes with homology 15 to the gene of interest (Ronchi et al 1995 EMBO J. 14: The other mechanism of co-suppression is 5318-5328). where the combined levels post-transcriptional, expression from both the gene and the transgene is thought to produce high levels of transcript which 20 triggers threshold-induced degradation of both messages (van Bokland et al., 1994 Plant J. 6: 861-877). exact molecular basis for co-suppression is unknown.

Unfortunately, both antisense and co-suppression technologies are subject to problems in heritability of the desired trait (Finnegan and McElroy 1994 Bio/Technology 12: 883-888). Currently, there is no easy way to specifically inactivate a gene of interest at the DNA level in plants (Pazkowski et al., 1988 EMBO J. 7: 4021-4026). Transposon mutagenesis is inefficient and not a stable event, while chemical mutagenesis is highly non-specific.

Applicant believes that ribozymes present an attractive alternative and because of their catalytic mechanism of action, have advantages over competing

technologies. However, there have been difficulties in effectiveness of ribozymes the demonstrating modulating gene expression in plant systems (Mazzolini et al., 1992 Plant Mol. Biol. 20: 715-731; Kull et al., 5 1995 J. Genet. & Breed. 49: 69-76). Although there are reports in the literature of ribozyme activity in plants cells, almost all of them involve down regulation of exogenously introduced genes, such as reporter genes in transient assays (Steinecke et al., 1992 EMBO J. 11:1525-1530; Perriman et al., 1993 Antisense Res. Dev. 3: 253-263; Perriman et al., 1995, Proc. Natl. Acad. Sci. USA, 92, 6165).

There are also several publications, [e.g., Lamb and Hay, 1990, J. Gen. Virol. 71, 2257-2264; Gerlach et al., 15 International PCT Publication No. WO 91/13994; Xu et al., 1992, Science in China (Ser. B) 35, 1434-1443; Edington 1992, in Gene Regulation: Biology of and Nelson, antisense RNA and DNA, eds. R. P. Erickson and J. G. Izant, pp 209-221, Raven Press, NY.; Atkins et al., 20 International PCT Publication No. WO 94/00012; Lenee et al., International PCT Publication Nos. WO 94/19476 and WO 9503404, Atkins et al., 1995, J. Gen. Virol. 76, 1781-1790; Gruber et al., 1994, <u>J. Cell. Biochem.</u> Suppl. 18A, 110 (X1-406) and Feyter et al., 1996, Mol. Gen. Genet. 25 250, 329-338], that propose using hammerhead ribozymes to modulate: virus replication, expression of viral genes and/or reporter genes. None of these publications report the use of ribozymes to modulate the expression of plant genes.

Mazzolini et al., 1992, Plant. Mol. Bio. 20, 715-731; Steinecke et al., 1992, EMBO. J. 11, 1525-1530; Perriman et al., 1995, Proc. Natl. Acad. Sci. USA., 92, 6175-6179; Wegener et al., 1994, Mol. Gen. Genet. 245, 465-470; and Steinecke et al., 1994, Gene, 149, 47-54,

describe the use of hammerhead ribozymes to inhibit expression of reporter genes in plant cells.

Bennett and Cullimore, 1992 Nucleic Acids Res. 20, 831-837 demonstrate hammerhead ribozyme-mediated in vitro cleavage of glna, glnb, glng and glnd RNA, coding for glutamine synthetase enzyme in Phaseolus vulgaris.

undesirable contain alkaloid Certain plants compounds which, when present in excess, are undesirable for human or animal consumption (Valkonen et al. 1996 10 Crit. Rev. Plant Sci. 15, 1-20). Potatoes and other solanaceous plants contain steroidal glycoalkaloids, whose regulated by genetic, developmental environmental signals (Bergenstrahle et al. 1992 J. Plant Phys. 140, 269-275; Sinden, 1984 Am. Potato J. 61, 141-156). Potato tubers synthesize the alkaloids solanine and chaconine in response to wounding, temperature, light and sprouting. These glycoalkaloids are thought to be responsible for preventing predation insect resistance to infection by pathogenic fungi (Valkonen et 20 al. supra). The enzyme solanidine UDP-glucose glucosyltransferase is implicated as the enzyme primarily responsible for the biosynthesis of both these alkaloid compounds (Stapleton et al. 1992 Prot. Exp. Purif. 3, 85-92, 6; Stapleton et al. 1991 J. Agri. Food Chem. 39, 25 1187-1193).

The mitochondrial tricarboxylic acid (TCA) cycle enzyme citrate synthase is implicated in the formation of flower buds in plants (Landshutze et al., 1995 EMBO J. 14, 660-666). Experiments with antisense constructs have shown that inhibition of the expression of the gene for this enzyme can delay or eliminate flower bud formation. There were no visible effects on plant growth or yield. The ovaries in the transgenic antisense plants disintegrated, indicating that citrate synthase and the

TCA cycle are important in the transition from vegetative to generative phase of plant growth. Cytoplasmic male sterility (CMS) has been associated with mitochondrial gene expression, but typically affects the ability of the 5 plant to produce viable pollen, not affecting female fertility (Levings et al., 1993 Plant Cell 5, 1285-1290; 5, 1277-1283). Inhibition Chaudhury, 1993 Plant Cell of expression of the citrate synthase gene by ribozymes should result in the delay or elimination of flower 10 formation in plants. This would be very useful in in plant species that preventing flowering vegetatively propagated or where the primary consumable part of the plant is root, stem or leaf. The enzyme is mitochondrial, but is encoded by a nuclear (Landshutze et al., 1995 Planta 196, 756-764). Chemical 15 mitochondrial respiration is inhibition of (Kromer et al., 1991 Plant. Phys. 95, 1270-1276), thus the ribozyme genetic approach is potentially advantageous over other methods.

The references cited above are distinct from the presently claimed invention since they do not disclose and/or contemplate the use of ribozymes to down regulate genes involved in the plant alkaloid biosynthesis in plant cells, let alone plants.

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Summary Of The Invention

The invention features modulation of gene expression in plants specifically using enzymatic nucleic acid molecules. Preferably, invention features inhibiting the expression of genes involved in the biosynthesis of certain alkaloid compounds using enzymatic nucleic acid molecules. That is, the inhibition of the gene product (e.g., RNA) results in a lowering of the production of alkaloid in the plant. Limiting the levels of certain

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alkaloid compounds in commercial cultivars, especially reductions in alkaloid content in the tuber by use of tissue-specific promoters is disclosed. The isolation of the gene encoding solanidine glucosyltransferase now allows evaluation of the phenotype that results from down-regulation of this gene (Moehs et al., 1997 Plant J. 11, 100-110). This application further deals with methods to produce cultivars such as, potato, tomato, pepper, eggplant, ditura, and others, with low levels of the toxic alkaloids.

In another aspect, the invention features inhibiting the expression of genes involved in flower formation using enzymatic nucleic acid molecules. That is, the inhibited to prevent gene product (e.g., RNA) is 15 formation of a flower by the plant modulating the expression of citrate synthase in commercial cultivars by use of enzymatic nucleic acid is disclosed as one example. Inhibition of expression of the synthase gene by ribozymes may result in the delay or 20 elimination of flower formation in plants. This would be very useful in preventing flowering in plant species that vegetatively propagated or where the consumable part of the plant is root, stem or leaf. This application further deals with methods to 25 cultivars such as, lettuce, spinach, cabbage, brussel sprouts, arugula, kale, collards, chard, beet, turnip, potato, sweet potato and turfgrass, with delayed or elimination of flower formation. Any gene in the flower formation pathway that does not effect vegetative growth 30 can be targeted in this manner.

The enzymatic nucleic acid molecule with RNA cleaving activity may be in the form of, but not limited to, a hammerhead, hairpin, hepatitis delta virus, group I intron, group II intron, RNaseP RNA, Neurospora VS RNA

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and the like. The enzymatic nucleic acid molecule with RNA cleaving activity may be encoded as a monomer or a multimer, preferably a multimer. The nucleic acids encoding for the enzymatic nucleic acid molecule with RNA 5 cleaving activity may be operably linked to an open reading frame. Gene expression in any plant species may be modified by transformation of the plant with the nucleic acid encoding the enzymatic nucleic acid molecules with RNA cleaving activity. There are also 10 numerous technologies for transforming a plant: such technologies include but are not limited to transformation with Agrobacterium, bombarding with DNA coated microprojectiles, whiskers, or electroporation. target gene may be modified with the nucleic acids 15 encoding the enzymatic nucleic acid molecules with RNA cleaving activity.

Ribozymes can be used to modulate flower formation of a plant, for example, by modulating the activity of an enzyme involved in a biochemical pathway. It may be desirable, in some instances, to decrease the level of expression of a particular gene, rather than shutting down expression completely: ribozymes can be used to achieve this. Enzymatic nucleic acid-based techniques were developed herein to allow directed modulation of gene expression to generate plant cells, plant tissues or plants with altered flowering phenotype.

In a preferred embodiment the invention features Ribozymes that can be used to modulate a specific trait of a plant cell, for example, by modulating the activity of an enzyme involved in a biochemical pathway. It may be desirable, in some instances, to decrease the level of expression of a particular gene, rather than shutting down expression completely: ribozymes can be used to achieve this. Enzymatic nucleic acid-based techniques

were developed herein to allow directed modulation of gene expression to generate plant cells, plant tissues or plants with altered phenotype.

Ribozymes (i.e., enzymatic nucleic acids) are nucleic acid molecules having an enzymatic activity which is able to repeatedly cleave other separate RNA molecules in a nucleotide base sequence-specific manner. Such enzymatic RNA molecules can be targeted to virtually any RNA transcript, and efficient cleavage has been achieved in vitro and in vivo (Zaug et al., 1986, Nature 324, 429; Kim et al., 1987, Proc. Natl. Acad. Sci. USA 84, 8788; Dreyfus, 1988, Einstein Quarterly J. Bio. Med., 6, 92; Haseloff and Gerlach, 1988, Nature 334 585; Cech, 1988, JAMA 260, 3030; Murphy and Cech, 1989, Proc. Natl. Acad. Sci. USA., 86, 9218; Jefferies et al., 1989, Nucleic Acids Research 17, 1371).

sequence-specificity, transof their Because cleaving ribozymes may be used as efficient tools to modulate gene expression in a variety of organisms 20 including plants, animals and humans (Bennett et al., supra; Edington et al., supra; Usman & McSwiggen, 1995 Ann. Rep. Med. Chem. 30, 285-294; Christoffersen and Marr, 1995 J. Med. Chem. 38, 2023-2037). Ribozymes can be designed to cleave specific RNA targets within the 25 background of cellular RNA. Such a cleavage event renders the mRNA non-functional and abrogates protein expression from that RNA. In this manner, synthesis of a protein associated with a particular phenotype and/or disease state can be selectively inhibited.

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

Brief Description of the Figures

Figure 1 is a diagrammatic representation of the hammerhead ribozyme domain known in the art. Stem II can be ≥ 2 base-pairs long. Each N is any nucleotide and 5 each • represents a base pair.

Figure 2a is a diagrammatic representation of the hammerhead ribozyme domain known in the art; Figure 2b is a diagrammatic representation of the hammerhead ribozyme as divided by Uhlenbeck (1987, Nature, 327, 596-600) into a substrate and enzyme portion; Figure 2c is a similar diagram showing the hammerhead divided by Haseloff and Gerlach (1988, Nature, 334, 585-591) into two portions; and Figure 2d is a similar diagram showing the hammerhead divided by Jeffries and Symons (1989, Nucl. Acids. Res., 17, 1371-1371) into two portions.

Figure 3 is a diagrammatic representation of the general structure of a hairpin ribozyme. Helix 2 (H2) is provided with a least 4 base pairs (i.e., n is 1, 2, 3 or 4) and helix 5 can be optionally provided of length 2 20 or more bases (preferably 3 - 20 bases, i.e., m is from 1 - 20 or more). Helix 2 and helix 5 may be covalently linked by one or more bases (i.e., r is \geq 1 base). Helix 1, 4 or 5 may also be extended by 2 or more base pairs (e.g., 4 - 20 base pairs) to stabilize the ribozyme 25 structure, and preferably is a protein binding site. each instance, each N and N' independently is any normal or modified base and each dash represents a potential These nucleotides may be base-pairing interaction. modified at the sugar, base or phosphate. Complete base-30 pairing is not required in the helices, but is preferred. Helix 1 and 4 can be of any size (i.e., o and p is each independently from 0 to any number, e.g., 20) as long as some base-pairing is maintained. Essential bases are

shown as specific bases in the structure, but those in the art will recognize that one or more may be modified and/or phosphate base, sugar chemically (abasic, modifications) or replaced with another base without Helix 4 can be formed from two 5 significant effect. separate molecules, i.e., without a connecting loop. connecting loop when present may be a ribonucleotide with or without modifications to its base, sugar or phosphate. "q" is \geq 2 bases. The connecting loop can also be 10 replaced with a non-nucleotide linker molecule. H refers to bases A, U, or C. Y refers to pyrimidine bases. " refers to a covalent bond.

Figure 4 is a representation of the general structure of the hepatitis Δ virus ribozyme domain known 15 in the art.

Figure 5 is a representation of the general structure of the self-cleaving VS RNA ribozyme domain.

Detailed Description Of The Invention

20 The present invention concerns compositions and methods for the modulation of gene expression in plants specifically using enzymatic nucleic acid molecules.

The following phrases and terms are defined below:

By "inhibit" or "modulate" is meant that the activity of enzymes, such as solanidine UDP-glucose glucosyl-transferase, potato citrate synthase, or level of mRNAs encoded by these genes is reduced below that observed in the absence of an enzymatic nucleic acid and preferably is below that level observed in the presence of an inactive RNA molecule able to bind to the same site on the mRNA, but unable to cleave that RNA.

By "enzymatic nucleic acid molecule" it is meant a nucleic acid molecule which has complementarity in a substrate binding region to a specified gene target, and also has an enzymatic activity which is active specifically cleave that target. That is, the enzymatic nucleic acid molecule is able to intermolecularly cleave DNA) and thereby inactivate a target RNA molecule. This complementarity functions to allow sufficient hybridization of the enzymatic nucleic acid 10 molecule to the target RNA to allow the cleavage to occur. One hundred percent complementarity is preferred, but complementarity as low as 50-75% may also be useful in this invention. The nucleic acids may be modified at the base, sugar, and/or phosphate groups. The term 15 enzymatic nucleic acid is used interchangeably with phrases such as ribozymes, catalytic RNA, enzymatic RNA, catalytic DNA, nucleozyme, DNAzyme, RNA enzyme, RNAzyme, polyribozymes, molecular scissors, self-splicing self-cleaving RNA, cis-cleaving RNA, autolytic RNA, 20 endoribonuclease, minizyme, leadzyme, oligozyme or DNA enzyme. All of these terminologies describe nucleic acid molecules with enzymatic activity. The term encompasses enzymatic RNA molecule which include one or ribonucleotides and may include a majority of other types 25 of nucleotides or abasic moieties, as described below.

By "complementarity" is meant a nucleic acid that can form hydrogen bond(s) with other RNA sequences by either traditional Watson-Crick or other non-traditional types (for example, Hoogsteen type) of base-paired interactions.

By "vectors" is meant any nucleic acid- and/or viral-based technique used to deliver and/or express a desired nucleic acid.

By "gene" is meant a nucleic acid that encodes an RNA.

By "plant gene" is meant a gene encoded by a plant.

By "endogenous" gene is meant a gene normally found in a plant cell in its natural location in the genome.

By "foreign" or "heterologous" gene is meant a gene not normally found in the host plant cell, but that is introduced by standard gene transfer techniques.

By "nucleic acid" is meant a molecule which can be 10 single-stranded or double-stranded, composed of nucleotides containing a sugar, a phosphate and either a purine or pyrimidine base which may be same or different, and may be modified or unmodified.

By "genome" is meant genetic material contained in 15 each cell of an organism and/or a virus.

By "mRNA" is meant RNA that can be translated into protein by a cell.

By "cDNA" is meant DNA that is complementary to and derived from a mRNA.

By "dsDNA" is meant a double stranded cDNA. 20

By "sense" RNA is meant RNA transcript that comprises the mRNA sequence.

By "antisense RNA" is meant an RNA transcript that comprises sequences complementary to all or part of a 25 target RNA and/or mRNA and that blocks the expression of a target gene by interfering with the processing, transport and/or translation of its primary transcript The complementarity may exist with any and/or mRNA. part of the target RNA, i.e., at the 5' non-coding 30 sequence, 3' non-coding sequence, introns, or the coding sequence. Antisense RNA is normally a mirror image of the sense RNA.

"expression", as used herein, is meant the transcription and stable accumulation of the enzymatic

nucleic acid molecules, mRNA and/or the antisense RNA Expression of genes involves inside a plant cell. transcription of the gene and translation of the mRNA into precursor or mature proteins.

By "cosuppression" is meant the expression of a foreign gene, which has substantial homology to an gene, and in a plant cell causes the reduction in activity in of the foreign and/or the endogenous protein product.

By "altered levels" is meant the level of production 10 of a gene product in a transgenic organism is different from that of a normal or non-transgenic organism.

By "promoter" is meant nucleotide sequence element within a gene which controls the expression of that gene. Promoter sequence provides the recognition for 15 polymerase and other transcription factors required for Promoters from a variety of efficient transcription. sources can be used efficiently in plant cells to express ribozymes. For example, promoters of bacterial origin, such as the octopine synthetase promoter, the nopaline 20 synthase promoter, the manopine synthetase promoter; promoters of viral origin, such as the cauliflower mosaic virus (35S); plant promoters, such as the ribulose-1,6biphosphate (RUBP) carboxylase small subunit (ssu), the beta-conglycinin promoter, the phaseolin promoter, ADH promoter, heat-shock promoters, and tissue specific Promoter may also contain certain enhancer promoters. sequence elements that may improve the transcription efficiency.

By "enhancer" is meant nucleotide sequence element which can stimulate promoter activity (Adh).

By "constitutive promoter" is meant promoter element that directs continuous gene expression in all cells types and at all times (actin, ubiquitin, CaMV 35S).

By "tissue-specific" promoter is meant promoter element responsible for gene expression in specific cell or tissue types, such as the leaves or seeds (zein, oleosin, napin, ACP).

By "development-specific" promoter is meant promoter element responsible for gene expression at specific plant developmental stage, such as in early or late embryogenesis.

By "inducible promoter" is meant promoter element which is responsible for expression of genes in response to a specific signal, such as: physical stimulus (heat shock genes); light (RUBP carboxylase); hormone (Em); metabolites; and stress.

By a "plant" is meant a photosynthetic organism, 15 either eukaryotic and prokaryotic.

By "angiosperm" is meant a plant having its seed enclosed in an ovary (e.g., coffee, tobacco, bean, pea).

By "gymnosperm" is meant a plant having its seed exposed and not enclosed in an ovary (e.g., pine, 20 spruce).

By "monocotyledon" is meant a plant characterized by the presence of only one seed leaf (primary leaf of the embryo). For example, maize, wheat, rice and others.

By "dicotyledon" is meant a plant producing seeds 25 with two cotyledons (primary leaf of the embryo). For example, coffee, canola, peas and others.

By "transgenic plant" is meant a plant expressing a foreign gene.

By "open reading frame" is meant a nucleotide 30 sequence, without introns, encoding an amino acid sequence, with a defined translation initiation and termination region.

The invention provides a method for producing a class of enzymatic cleaving agents which exhibit a high

degree of specificity for the RNA of a desired target.

The enzymatic nucleic acid molecule may be targeted to a highly specific sequence region of a target such that specific gene inhibition can be achieved. Alternatively, enzymatic nucleic acid can be targeted to a highly conserved region of a gene family to inhibit gene expression of a family of related enzymes. The ribozymes can be expressed in plants that have been transformed with vectors which express the nucleic acid of the present invention.

The enzymatic nature of a ribozyme is advantageous over other technologies, since the concentration ribozyme necessary to affect a therapeutic treatment is This advantage reflects the ability of the lower. 15 ribozyme to act enzymatically. Thus, a single ribozyme molecule is able to cleave many molecules of target RNA. is a highly specific ribozyme addition, the inhibitor, with the specificity of inhibition depending not only on the base-pairing mechanism of binding to the 20 target RNA, but also on the mechanism of target RNA cleavage. Single mismatches, or base-substitutions, near the site of cleavage can completely eliminate catalytic activity of a ribozyme.

enzymatic RNAs are known presently. Each can catalyze the hydrolysis of RNA phosphodiester bonds in trans (and thus can cleave other RNA molecules) under physiological conditions. Table I summarizes some of the characteristics of these ribozymes. In general, enzymatic nucleic acids act by first binding to a target RNA. Such binding occurs through the target binding portion of an enzymatic nucleic acid which is held in close proximity to an enzymatic portion of the molecule that acts to cleave the target RNA. Thus, the enzymatic nucleic acid first

recognizes and then binds a target RNA through complementary base-pairing, and once bound to the correct site, acts enzymatically to cut the target RNA. Strategic cleavage of such a target RNA will destroy its ability to direct synthesis of an encoded protein. After an enzymatic nucleic acid has bound and cleaved its RNA target, it is released from that RNA to search for another target and can repeatedly bind and cleave new targets.

the preferred embodiments one of 10 In inventions herein, the enzymatic nucleic acid molecule is formed in a hammerhead or hairpin motif, but may also be formed in the motif of a hepatitis Δ virus, group I intron, group II intron or RNaseP RNA (in association 15 with an RNA guide sequence) or Neurospora VS RNA. Examples of such hammerhead motifs are described by Dreyfus, supra, Rossi et al., 1992, AIDS Research and Human Retroviruses 8, 183; of hairpin motifs by Hampel et al., EP0360257, Hampel and Tritz, 1989 Biochemistry 20 4929, Feldstein et al., 1989, Gene 82, 53, Haseloff and Gerlach, 1989, Gene, 82, 43, and Hampel et al., 1990 Nucleic Acids Res. 18, 299; of the hepatitis Δ virus described by Perrotta and Been, is motif Biochemistry 31, 16; of the RNaseP motif by Guerrier-Takada et al., 1983 Cell 35, 849; Forster and Altman, 1990, <u>Science</u> 249, 783; Li and Altman, 1996, <u>Nucleic</u> Acids Res. 24, 835; Neurospora VS RNA ribozyme motif is described by Collins (Saville and Collins, 1990 Cell 61, 685-696; Saville and Collins, 1991 Proc. Natl. Acad. Sci. USA 88, 8826-8830; Collins and Olive, 1993 Biochemistry 32, 2795-2799; Guo and Collins, 1995, EMBO. J. 14, 363); Group II introns are described by Griffin et al., 1995, Chem. Biol. 2, 761; Michels and Pyle, 1995, Biochemistry

34, 2965; and of the Group I intron by Cech et al., U.S. Patent 4,987,071. These specific motifs are not limiting in the invention and those skilled in the art will recognize that all that is important in an enzymatic nucleic acid molecule of this invention is that it has a specific substrate binding site which is complementary to one or more of the target gene RNA regions, and that it have nucleotide sequences within or surrounding that substrate binding site which impart an RNA cleaving activity to the molecule.

The enzymatic nucleic acid molecules of the instant invention will be expressed within cells from eukaryotic promoters [e.g., Gerlach et al., International Publication No. WO 91/13994; Edington and Nelson, 1992, 15 in Gene Regulation: Biology of Antisense RNA and DNA, eds. R. P. Erickson and J. G. Izant, pp 209-221, Raven Press, NY.; Atkins et al., International PCT Publication International No. WO 94/00012; Lenee et al., Publication Nos. WO 94/19476 and WO 9503404, Atkins et 20 al., 1995, J. Gen. Virol. 76, 1781-1790; McElroy and Brettell, 1994, <u>TIBTECH</u> 12, 62; Gruber <u>et al.</u>, 1994, <u>J.</u> Cell. Biochem. Suppl. 18A, 110 (X1-406) and Feyter et al., 1996, Mol. Gen. Genet. 250, 329-338; all of these are incorporated by reference herein]. Those skilled in the art will realize from the teachings herein that any 25 ribozyme can be expressed in eukaryotic plant cells from an appropriate promoter. The ribozymes expression is under the control of a constitutive promoter, a tissuespecific promoter or an inducible promoter.

To obtain the ribozyme mediated modulation, the ribozyme RNA is introduced into the plant. There are also numerous ways to transform plants; plants can be transformed using the gene gun (US Patents 4,945,050 to Cornell and 5,141,131 to DowElanco); plants may be

transformed using Agrobacterium technology, see US Patent 5,177,010 to University of Toledo, 5,104,310 to Texas A&M, European Patent Application 0131624B1, Patent Applications 120516, 159418B1 and 176,112 5 Schilperoot, US Patents 5,149,645, 5,469,976, 5,464,763 and 4,940,838 and 4,693,976 to Schilperoot, Applications 116718, 290799, 320500 all Patent Applications 604662 European MaxPlanck, 627752 to Japan Tobacco, European Patent Applications 10 0267159, and 0292435 and US Patent 5,231,019 all to Ciba Patents 5,463,174 and 4,762,785 both to US Calgene, and US Patents 5,004,863 and 5,159,135 both to Agracetus; whiskers technology, see US Patents 5,302,523 and 5,464,765 both to Zeneca; electroporation technology, 15 see WO 87/06614 to Boyce Thompson Institute, 5,472,869 and 5,384,253 both to Dekalb, WO9209696 and WO9321335 both to PGS; all of which are incorporated by reference herein in totality. In addition to numerous technologies for transforming plants, the type of tissue which is contacted with the foreign material (typically plasmids 20 containing RNA or DNA) may vary as well. Such tissue would include but would not be limited to embryogenic and any tissue tissue, callus tissue type I and II, which is receptive to transformation and subsequent 25 regeneration into a transgenic plant. Another variable is the choice of a selectable marker. The preference for a particular marker is at the discretion of the artisan, but any of the following selectable markers may be used along with any other gene not listed herein which could function as a selectable marker. Such selectable markers chlorosulfuron, not limited to but are include hygromyacin, PAT and/or bar, bromoxynil, kanamycin and the like. The bar gene may be isolated from Strptomuces, particularly from the hygroscopicus or viridochromogenes species. The bar gene codes for phosphinothricin acetyl transferase (PAT) that inactivates the active ingradient in the herbicide bialaphos phosphinothricin (PPT). Thus, numerous combinations of technologies may be used in employing ribozyme mediated modulation.

The ribozymes may be expressed individually monomers, i.e., one ribozyme targeted against one site is expressed per transcript. Alternatively, two or more ribozymes targeted against more than one target site are 10 expressed as part of a single RNA transcript. RNA transcript comprising more than one ribozyme targeted against more than one cleavage site are readily generated to achieve efficient modulation of gene expression. Ribozymes within these multimer constructs are the same 15 or different. For example, the multimer construct may comprise a plurality of hammerhead ribozymes or hairpin ribozymes or other ribozyme motifs. Alternatively, the multimer construct may be designed to include a plurality of different ribozyme motifs, such as hammerhead and 20 hairpin ribozymes. More specifically, multimer ribozyme constructs are designed, wherein a series of ribozyme motifs are linked together in tandem in a single RNA The ribozymes are linked to each other by transcript. nucleotide linker sequence, wherein the linker sequence 25 may or may not be complementary to the target RNA. Multimer ribozyme constructs (polyribozymes) are likely effectiveness of ribozyme-mediated the improve modulation of gene expression.

The activity of ribozymes can also be augmented by their release from the primary transcript by a second ribozyme (Draper et al., PCT WO 93/23569, and Sullivan et al., PCT WO 94/02595, both hereby incorporated in their totality by reference herein; Ohkawa, J., et al., 1992, Nucleic Acids Symp. Ser., 27, 15-6; Taira, K., et al.,

1991, <u>Nucleic Acids Res.</u>, 19, 5125-30; Ventura, M., <u>et al.</u>, 1993, <u>Nucleic Acids Res.</u>, 21, 3249-55; Chowrira <u>et al.</u>, 1994 <u>J. Biol. Chem.</u> 269, 25856).

Ribozyme-mediated modulation of gene expression can be practiced in a wide variety of plants including but not limited to potato, lettuce spinach, cabbage, brussel sprouts, arugula, kale, collards, chard, beet, turnip, sweet potato and turfgrass. Following are a few non-limiting examples that describe the general utility of ribozymes in modulation of gene expression.

Thus, in one instance, the invention concerns compositions (and methods for their for use) modulation of genes involved in the biosynthesis of undesirable alkaloid compounds in plants. This is 15 accomplished through the inhibition of genetic expression, with ribozymes, which results in the reduction or elimination of certain gene activities in plants, such as glucosyl-transferase. solanidine UDP-glucose activity is reduced in plants, such as potato and other These endogenously expressed 20 solanaceous plants. ribozyme molecules contain substrate binding domains that bind to accessible regions of the target RNA. molecules also contain domains that catalyze the cleavage The RNA molecules are preferably ribozymes of of RNA. Upon binding, the 25 the hammerhead or hairpin motif. ribozymes cleave the target mRNAs, preventing translation In the absence of and protein accumulation. expression of the target gene, and/or if the level of expression of the target gene is significantly reduced, 30 levels of undesirable alkaloids is reduced or inhibited. Specific examples are provided below in the Tables III and IV.

In one aspect, the ribozymes have binding arms which are complementary to the substrate sequences in Tables

III and IV. Those in the art will recognize that while such examples are designed to one gene RNA (solanidine UDP-glucose glucosyl-transferase) of one plant(e.g., potato), similar ribozymes can be made complementary to other genes in other plant's RNA. By complementary is thus meant that the binding arms of the ribozymes are able to interact with the target RNA in a sequencespecific manner and enable the ribozyme to cause cleavage of a plant mRNA target. Examples of such ribozymes are 10 typically sequences defined in Tables III and IV. The active ribozyme typically contains an enzymatic center equivalent to those in the examples, and binding arms able to bind plant mRNA such that cleavage at the target site occurs. Other sequences may be present which do not 15 interfere with such binding and/or cleavage.

features instance, invention another the In (and methods for their use) for the compositions modulation of genes involved in the flower formation in plants. This is accomplished through the inhibition of 20 genetic expression, with ribozymes, which results in the reduction or elimination of certain gene activities in plants, such as citrate synthase. Such activity can be reduced in plants, such as lettuce, spinach, cabbage, brussel sprouts, arugula, kale, collards, chard, beet, turnip, potato, sweet potato and turfgrass. These endogenously expressed ribozyme molecules contain substrate binding domains that bind to accessible regions of the The RNA molecules also contain domains that target RNA. The RNA molecules are catalyze the cleavage of RNA. 30 preferably ribozymes of the hammerhead or hairpin motif. Upon binding, the ribozymes cleave the target mRNAs, preventing translation and protein accumulation. absence of the expression of the target gene, and/or if level of expression of the target gene is the

significantly reduced, levels of undesirable alkaloids is Specific examples are provided reduced or inhibited. below in the Tables V and VI. In a non-limiting example, ribozymes have binding arms which are complementary to 5 the substrate sequences shown in Tables V and VI are disclosed. Those in the art will recognize that while such examples are designed to one gene RNA (citrate synthase) of one plant (e.g., potato), similar ribozymes can be made complementary to other genes in other plant's 10 RNA. By complementary is thus meant that the binding arms of the ribozymes are able to interact with the target RNA in a sequence-specific manner and enable the ribozyme to cause cleavage of a plant mRNA target. Examples of such ribozymes are typically sequences 15 defined in Tables V and VI. The active ribozyme typically contains an enzymatic center equivalent to those in the examples, and binding arms able to bind plant mRNA such that cleavage at the target site occurs. Other sequences may be present which do not interfere with such binding 20 and/or cleavage.

The sequences of the ribozymes that are particularly useful in this study, are shown in Tables III-VI.

Those in the art will recognize that ribozyme sequences listed in the Tables are representative only of 25 many more such sequences where the enzymatic portion of the ribozyme (all but the binding arms) is altered to affect activity. For example, stem-loop II sequence of hammerhead ribozymes listed in Table III and V (5'-GGCGAAAGCC-3') can be altered (substitution, deletion, and/or insertion) to contain any sequences, preferably provided that a minimum of a two base-paired stem structure can form. Similarly, stem-loop IV sequence of hairpin ribozymes listed in Table IV and VI (5'-CACGUUGUG-3') can be altered (substitution, deletion,

and/or insertion) to contain any sequence, preferably provided that a minimum of a two base-paired stem structure can form. Such ribozymes are equivalent to the ribozymes described specifically in the Tables.

Preferably, the recombinant vectors capable of stable integration into the plant genome and selection of transformed plant lines expressing the ribozymes are expressed either by constitutive or inducible promoters in the plant cells. Once expressed, the ribozymes cleave their target mRNAs and reduce alkaloid production in their host cells. The ribozymes expressed in plant cells are under the control of a constitutive promoter, a tissue-specific promoter or an inducible promoter.

Modification of undesirable alkaloid profile is an important application of nucleic acid-based technologies which are capable of reducing specific gene expression. A high level of undesirable alkaloid compounds is undesirable in plants that produce products of commercial importance.

In preferred embodiments, hairpin and hammerhead ribozymes that cleave solanidine UDP-glucose glucosyltransferase RNA are described. Those of ordinary skill in the art will understand from the examples described below that other ribozymes that cleave target RNAs required for solanidine UDP-glucose glucosyl-transferase activity may now be readily designed and are within the scope of the invention.

Modification of flower formation is an important application of nucleic acid-based technologies which are capable of reducing specific gene expression. In preferred embodiments, hairpin and hammerhead ribozymes that cleave potato citrate synthase RNA are described. Those of ordinary skill in the art will understand from the examples described below that other ribozymes that

cleave target RNAs required for potato citrate synthase activity may now be readily designed and are within the scope of the invention

While specific examples to potato RNA are provided,

those in the art will recognize that the teachings are
not limited to potato. Furthermore, the same or
equivalent target may be used in other plant species.
The complementary arms suitable for targeting the
specific plant RNA sequences are utilized in the ribozyme
targeted to that specific RNA. The examples and teachings
herein are meant to be non-limiting, and those skilled in
the art will recognize that similar embodiments can be
readily generated in a variety of different plants to
modulate expression of a variety of different genes,
using the teachings herein, and are within the scope of
the inventions.

Standard molecular biology techniques were followed in the examples herein. Additional information may be found in Sambrook, J., Fritsch, E. F., and Maniatis, T. (1989), Molecular Cloning a Laboratory Manual, second edition, Cold Spring Harbor: Cold Spring Harbor Laboratory Press, which is incorporated herein by reference.

25 Examples

30

Example 1: Identification of Potential Ribozyme Cleavage Sites for solanidine UDP-glucose glucosyl-transferase

Approximately 353 HH ribozyme cleavage sites and approximately 20 HP sites were identified in the potato solanidine UDP-glucose glucosyl-transferase RNA. A HH site consists of a uridine and any nucleotide except guanosine (UH). Tables III and IV have a list of HH and HP ribozyme cleavage sites. The numbering system starts with 1 at the 5' end of a solanidine UDP-glucose

glucosyl-transferase RNA having the sequence shown in Moehs et al., supra.

Ribozymes, such as those listed in Tables III and IV, can be readily designed and synthesized to such cleavage sites with between 5 and 100 or more bases as substrate binding arms (see Figs. 1 - 5). These substrate binding arms within a ribozyme allow the ribozyme to interact with their target in a sequence-specific manner.

10

Example 2: Selection of Ribozyme Cleavage Sites for solanidine UDP-glucose glucosyl-transferase

The secondary structure of solanidine UDP-glucose glucosyl-transferase RNA was assessed by computer analysis using algorithms, such as those developed by M. Zuker (Zuker, M., 1989 Science, 244, 48-52). Regions of the mRNA that did not form secondary folding structures with RNA/RNA stems of over eight nucleotides and contained potential hammerhead ribozyme cleavage sites were identified.

Example 3: Hammerhead and Hairpin Ribozymes for solanidine UDP-glucose glucosyl-transferase

Hammerhead (HH) and hairpin (HP) ribozymes are subjected to analysis by computer folding and the ribozymes that had significant secondary structure are rejected.

The ribozymes are chemically synthesized. The general procedures for RNA synthesis have been described previously (Usman et al., 1987, J. Am. Chem. Soc., 109, 7845-7854 and in Scaringe et al., 1990, Nucl. Acids Res., 18, 5433-5341; Wincott et al., 1995, Nucleic Acids Res. 23, 2677). Small scale syntheses are conducted on a 394 Applied Biosystems, Inc. synthesizer using a modified 2.5

 μmol scale protocol with a 5 min coupling step for alkylsilyl protected nucleotides and 2.5 min coupling step for 2'-0-methylated nucleotides. Table II outlines the amounts, and the contact times, of the reagents used 5 in the synthesis cycle. A 6.5-fold excess (163 μL of 0.1 $M = 16.3 \mu mol)$ of phosphoramidite and a 24-fold excess of S-ethyl tetrazole (238 μL of 0.25 M = 59.5 μ mol) relative to polymer-bound 5'-hydroxyl was used in each coupling cycle. Average coupling yields on the 394, determined by 10 colorimetric quantitation of the trityl fractions, was 97.5-99%. Other oligonucleotide synthesis reagents for the 394: Detritylation solution was 2% TCA in methylene chloride (ABI); capping was performed with 16% $\underline{\mathrm{N}}\text{-Methyl}$ imidazole in THF (ABI) and 10% acetic anhydride/10% 2,6lutidine in THF (ABI); oxidation solution is 16.9 mM I_2 , 49 mM pyridine, 9% water in THF (Millipore). Synthesis Grade acetonitrile is used directly from the reagent bottle. \underline{S} -Ethyl tetrazole solution (0.25 M in acetonitrile) was made up from the solid obtained from 20 American International Chemical, Inc.

Deprotection of the RNA is performed as follows. The oligoribonucleotide, trityl-off, polymer-bound transferred from the synthesis column to a 4 mL glass screw top vial and suspended in a solution of methylamine (MA) at 65°C for 10 min. After cooling to -20°C, the supernatant is removed from the polymer support. with 1.0 mL times three support is washed EtOH: MeCN: H2O/3:1:1, vortexed and the supernatant is then added to the first supernatant. The combined super-30 natants, containing the oligoribonucleotide, are dried to a white powder.

oligoribonucleotide base-deprotected The resuspended in anhydrous TEA•HF/NMP solution (250 μL of a

solution of 1.5 mL N-methylpyrrolidinone, 750 µL TEA and 1.0 mL TEA·3HF to provide a 1.4 M HF concentration) and heated to 65°C for 1.5 h. The resulting, fully deprotected, oligomer is quenched with 50 mM TEAB (9 mL) prior to anion exchange desalting.

For anion exchange desalting of the deprotected oligomer, the TEAB solution is loaded onto a Qiagen 500 anion exchange cartridge (Qiagen Inc.) that is prewashed with 50 mM TEAB (10 mL). After washing the loaded cartridge with 50 mM TEAB (10 mL), the RNA is eluted with 2 M TEAB (10 mL) and dried down to a white powder.

Inactive hammerhead ribozymes are synthesized by substituting a U for G_5 and a U for A_{14} (numbering from (Hertel, K. J., et al., 1992, Nucleic Acids Res., 20, 3252).

The hairpin ribozymes are synthesized as described above for the hammerhead RNAs.

DNA Ribozymes can also synthesized be from templates using bacteriophage T7 RNA polymerase (Milligan Enzymol. 180, Uhlenbeck, 1989, Methods 20 and Ribozymes are purified by gel electrophoresis using general methods or are purified by high pressure liquid chromatography (HPLC; See Wincott et al., 1996, supra, the totality of which is hereby incorporated herein by 25 reference) and were resuspended in water. The sequences of the chemically synthesized ribozymes used in this study are shown below in Tables III and IV.

Example 4: Construction of Ribozyme expressing transcription units for solanidine UDP-glucose glucosyltransferase

Ribozymes targeted to cleave solanidine UDP-glucose glucosyl-transferase RNA can be endogenously expressed in

plants, either from genes inserted into the plant genome (stable transformation) or from episomal transcription units (transient expression) which are part of plasmid vectors or viral sequences. These ribozymes can be expressed via RNA polymerase I, II, or III plant or plant virus promoters (such as CaMV). Promoters can be either constitutive, tissue specific, or developmentally expressed.

10 Example 5: Identification of Potential Ribozyme Cleavage Sites for potato citrate synthase

Approximately 398 HH ribozyme cleavage sites and approximately 25 HP sites were identified in the potato citrate synthase RNA. A HH site consists of a uridine and any nucleotide except guanosine (UH). Tables V and VI have a list of HH and HP ribozyme cleavage sites.

Ribozymes, such as those listed in Tables III and IV, can be readily designed and synthesized to such cleavage sites with between 5 and 100 or more bases as substrate binding arms (see Figs. 1 - 5). These substrate binding arms within a ribozyme allow the ribozyme to interact with their target in a sequence-specific manner.

25 Example 6: Selection of Ribozyme Cleavage Sites for potato citrate synthase

The secondary structure of potato citrate synthase RNA was assessed by computer analysis using algorithms, such as those developed by M. Zuker (Zuker, M., 1989)

30 Science, 244, 48-52). Regions of the mRNA that did not form secondary folding structures with RNA/RNA stems of over eight nucleotides and contained potential hammerhead ribozyme cleavage sites were identified.

Example 7: Hammerhead and Hairpin Ribozymes for potato citrate synthase

Hammerhead (HH) and hairpin (HP) ribozymes are subjected to analysis by computer folding and the ribozymes that had significant secondary structure are rejected.

The ribozymes are synthesized as described above. The sequences of the chemically synthesized ribozymes used in this study are shown below in Tables V and VI.

10

Example 8: Construction of Ribozyme expressing transcription units for potato citrate synthase

Ribozymes targeted to cleave potato citrate synthase RNA can be endogenously expressed in plants, either from genes inserted into the plant genome (stable transformation) or from episomal transcription units (transient expression) which are part of plasmid vectors or viral sequences. These ribozymes can be expressed via RNA polymerase I, II, or III plant or plant virus promoters (such as CaMV). Promoters can be either constitutive, tissue specific, or developmentally expressed.

Example 9: Plant Transformation and Construction

25

There are several methods to genetically engineer plants (for a review see Gasser et al., 1989 Science 244, 1293-1299; Potrykus, 1991 Annu. Rev. Plant Physiol. Plant Mol. Biol. 42, 205-225; Gasser and Fraley, 1992 Scientific American June 1992 pp 62-69). These methods can be used to introduce the above ribozymes directly or via exression vectors. These methods include the following:

Helium blasting involves accelerating suspended DNAcoated gold particles towards and into prepared tissue targets. The device used was an earlier prototype to the one described in a DowElanco U.S. Patent (#5,141,131) which is incorporated herein by reference, although both function in a similar manner. The device consists of a high pressure helium source, a syringe containing the pneumatically-operated DNA/gold suspension, and а multipurpose valve which provides controlled linkage 10 between the helium source and a loop of pre-loaded DNA/gold suspension. Prior to blasting, tissue targets are covered with a sterile 104 micron stainless steel screen, which holds the tissue in place during impact. Next, targets are placed under vacuum in the main chamber The DNA-coated gold particles are 15 of the device. accelerated at the target 4 times using a helium pressure Each blast delivered 20 μl of DNA/gold of 1500 psi. suspension. Immediately post-blasting, the targets are placed back on maintenance medium plus osmoticum for a 16 20 to 24 hour recovery period.

Bombardment-mediated transformation Particle (Gordon-Kamm et al., 1990 The Plant Cell 2, 603-618; Potrykus, 1991 Annu. Rev. Plant Physiol. Plant Mol. Biol. 42, 205-225; Gasser and Fraley, 1992 Scientific American June 1992 pp 62-69; Vain et al., 1993 Plant Cell Rep. 12, 84-88; Weymann et al., 1993 In Vitro Cell. Dev. Biol. 29P, 33-37): This strategy involves bombardment of plant cells with minute (1-2 microns in diameter) metal particles (for example tungsten or gold particles) using 30 a "gene" gun (also referred to as "Biolistics" or The metal particles, coated with "particle" gun). ribozyme (ribozyme or genetic material plasmids), can penetrate the cell wall, without causing any irreversible damage to the cell, and deliver the genetic material to the cytoplasm.

Electroporation-mediated transformation (Fromm et al., 1986 Nature 319, 791-793; Rhodes et al., 1988 5 Science 240, 204-207; Potrykus, 1991 Annu. Rev. Plant Physiol. Plant Mol. Biol. 42, 205-225; Gasser and Fraley, 1992 Scientific American June 1992 pp 62-69; D'Halluin et al., 1992 The Plant Cell 4, 1495-1505; Sukhapinda et al., 1993 Plant Cell Rep. 13, 63-68; Laursen et al., 1994 10 Plant Mol. Biol. 24, 51-61): This technique involves permeabilizing the target cell membrane by using short high voltage electric pulses. Nucleic acids (ribozyme encoding plasmids) can pass through a permeabilized cell membrane and potentially integrate into the host genome 15 resulting in a transformed phenotype. Electroporation can be carried out on (a) plant protoplasts, plant cells lacking a cell wall, (Fromm et al., 1986 Nature 319, 791-793; Rhodes et al., 1988 Science 240, 204-207; Sukhapinda et al., 1993 Plant Cell Rep. 13, 63-68); (b) cultured 20 cells (Laursen et al., 1994 Plant Mol. Biol. 24, 51-61); (c) Plant tissue (D'Halluin et al., 1992 The Plant Cell 4, 1495-1505).

Agrobacterium-mediated transformation: This method uses a disarmed (disease causing genes are deleted)

25 species of Agrobacterium tumefaciens or Agrobacterium rizogenes (Potrykus, 1991 Annu. Rev. Plant Physiol. Plant Mol. Biol. 42, 205-225; Gasser and Fraley, 1992 Scientific American June 1992 pp 62-69). This organism transfers part of its DNA into plant cells (T-DNA).

Agrobacterium containing the recombinant T-DNA can be generated. Agrobacterium will infect and release the recombinant T-DNA into maize cells. The integration of

T-DNA into host DNA will result in a transformed phenotype.

Other Uses:

Potential usefulness of sequence-specific enzymatic nucleic acid molecules of the instant invention might have many of the same applications for the study of RNA that DNA restriction endonucleases have for the study of DNA (Nathans, D. and Smith, H. O., (1975) Ann. Rev. Biochem. 44:273). For example, the pattern of restriction fragments could be used to establish sequence relationships between two related plant RNAs, and large plant RNAs could be specifically cleaved to fragments of a size more useful for study. The ability to engineer sequence specificity of the ribozyme is ideal for cleavage of RNAs of unknown sequence.

Ribozymes of this invention may be used as tools to examine genetic drift and mutations within plant cells. The close relationship between ribozyme activity and the 20 structure of the target RNA allows the detection of mutations in any region of the molecule which alters the base-pairing and three-dimensional structure of target RNA. By using multiple ribozymes described in this invention, one may map nucleotide changes which are important to RNA structure and function in vitro, as well as in cells and tissues. Cleavage of target RNAs with ribozymes may be used to inhibit gene expression and define the role (essentially) of specified gene products in the synthesis of undesirable alkaloids in plants. 30 this manner, other genetic targets may be defined as important mediators of alkaloid production. experiments will lead to better modifications of the alkaloid production by affording the possibility of combinational concepts (e.g., multiple ribozymes targeted to different genes intermittent treatment with combinations of ribozymes and/or other chemical or biological molecules). Other in vitro uses of ribozymes of this invention are well known in the art, and include detection of the presence of mRNA associated with undesirable alkaloid production condition. Such RNA is detected by determining the presence of a cleavage product after treatment with a ribozyme using standard methodology.

Other embodiments are within the following claims.

Table I

Table I:

Characteristics of naturally occurring ribozymes

Group I Introns

- 5 o Size: ~150 to >1000 nucleotides.
 - Requires a U in the target sequence immediately 5' of the cleavage site.
 - Binds 4-6 nucleotides at the 5'-side of the cleavage site.
- 10 Reaction mechanism: attack by the 3'-011 of guanosine to generate cleavage products with 3'-OH and 5'- quanosine.
 - Additional protein cofactors required in some cases to help folding and maintainance of the active structure
- 15 [¹].
 - Over 300 known members of this class. Found as an intervening sequence in Tetrahymena thermophila rRNA, fungal mitochondria, chloroplasts, phage T4, bluegreen algae, and others.
- 20 Major structural features largely established through phylogenetic comparisons, mutagenesis, and biochemical studies [2,3].
 - Complete kinetic framework established for one ribozyme [4,5,6,7]
- 25 Studies of ribozyme folding and substrate docking underway [8,9,10].
 - \circ Chemical modification investigation of important residues well established [11,12].
- o The small (4-6 nt) binding site may make this ribozyme too non-specific for targeted RNA cleavage, however, the Tetrahymena group I intron has been used to repair a "defective" ß-galactosidase message by the ligation

Table I

of new ß-galactosidase sequences onto the defective message $[^{13}]$.

RNAse P RNA (M1 RNA)

- 5 o Size: ~290 to 400 nucleotides.
 - RNA portion of a ubiquitous ribonucleoprotein enzyme.
 - \circ Cleaves tRNA precursors to form mature tRNA [14].
 - \circ Reaction mechanism: possible attack by M²⁺-OH to generate cleavage products with 3'OH and 5'-phosphate.
- 10 RNAse P is found throughout the prokaryotes and eukaryotes. The RNA subunit has been sequenced from bacteria, yeast, rodents, and primates.
 - Recruitment of endogenous RNAse P for therapeutic applications is possible through hybridization of an
- 15 External Guide Sequence (EGS) to the target RNA
 - Important phosphate and 2' OH contacts recently identified [17,18]

Group II Introns

- 20 o Size: >1000 nucleotides.
 - Trans cleavage of target RNAs recently demonstrated
 19,20].
 - Sequence requirements not fully determined.
 - Reaction mechanism: 2'-OH of an internal adenosine
- generates cleavage products with 3'-OH and a "lariat" RNA containing a 3'-5' and a 2'-5' branch point.
 - o Only natural ribozyme with demonstrated participation in DNA cleavage $[^{21},^{22}]$ in addition to RNA cleavage and ligation.
- 30 \circ Major structural features largely established through phylogenetic comparisons [23].

Table I

- \circ Important 2' OH contacts beginning to be identified $\lceil 24 \rceil$
- Kinetic framework under development [25]

5 Neurospora VS RNA

- o Size: ~144 nucleotides.
- \circ Trans cleavage of hairpin target RNAs recently demonstrated $[^{26}]$
- Sequence requirements not fully determined.
- 10 Reaction mechanism: attack by 2'-OH 5' to the scissile bond to generate cleavage products with 2',3'-cyclic phosphate and 5'-OH ends.
 - Binding sites and structural requirements not fully determined.
- o Only 1 known member of this class. Found in Neurospora VS RNA.

Hammerhead Ribozyme

(see text for references)

- 20 · Size: ~13 to 40 nucleotides.
 - Requires the target sequence UH immediately 5' of the cleavage site.
 - Binds a variable number nucleotides on both sides of the cleavage site.
- 25 Reaction mechanism: attack by 2'-OH 5' to the scissile bond to generate cleavage products with 2',3'-cyclic phosphate and 5'-OH ends.
 - 14 known members of this class. Found in a number of
 plant pathogens (virusoids) that use RNA as the
- 30 infectious agent.
 - Essential structural features largely defined, including 2 crystal structures []

- Minimal ligation activity demonstrated (for engineering through in vitro selection) []
- Complete kinetic framework established for two or more ribozymes [].
- 5 Chemical modification investigation of important residues well established [].

Hairpin Ribozyme

- Size: ~50 nucleotides.
- 10 Requires the target sequence GUC immediately 3' of the cleavage site.
 - Binds 4-6 nucleotides at the 5'-side of the cleavage site and a variable number to the 3'- side of the cleavage site.
- o Reaction mechanism: attack by 2'-OH 5' to the scissile bond to generate cleavage products with 2',3'-cyclic phosphate and 5'-OH ends.
 - o 3 known members of this class. Found in three plant pathogen (satellite RNAs of the tobacco ringspot
- virus, arabis mosaic virus and chicory yellow mottle virus) which uses RNA as the infectious agent.
 - Essential structural features largely defined
 [27,28,29,30]
 - Ligation activity (in addition to cleavage activity)
- makes ribozyme amenable to engineering through in vitro selection [31]
 - Complete kinetic framework established for one ribozyme [31].
- Chemical modification investigation of important
 residues begun [33,34]

15

Table I

Hepatitis Delta Virus (HDV) Ribozyme

- o Size: ~60 nucleotides.
- \circ Trans cleavage of target RNAs demonstrated [31].
- Binding sites and structural requirements not fully
- determined, although no sequences 5' of cleavage site are required. Folded ribozyme contains a pseudoknot structure [36].
 - Reaction mechanism: attack by 2'-OH 5' to the scissile bond to generate cleavage products with 2,3'-cyclic phosphate and 5'-OH ends.
 - Only 2 known members of this class. Found in human HDV.
 - \circ Circular form of HDV is active and shows increased nuclease stability $[^{37}]$
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10 Table II: 2.5 µmol RNA Synthesis Cycle

| | Reagent | Equivalents | Amount | Wait Time* |
|----|--------------------|-------------|---------|---------------|
| 15 | Phosphoramidites | 6.5 | 163 µL | 2.5 |
| | S-Ethyl Tetrazole | 23.8 | 238 µL | 2.5 |
| | Acetic Anhydride | 100 | 233 µL | 5 sec |
| | N-Methyl Imidazole | 186 | 233 μL | 5 sec |
| | TCA | 83.2 | 1.73 mL | 21 sec |
| 20 | Iodine | 8.0 | 1.18 mL | 45 sec |
| | Acetonitrile | NA | 6.67 mL | NA |

^{*} Wait time does not include contact time during delivery.

Table III

Table III: Solanidine glucosyltransferase Hammerhead Ribozyme and Target Sequences

| Nt. | Substrate | Ribozyme |
|----------|--------------------|---------------------------------|
| Position | | - CAR ROCCARCA |
| 13 | UCUUGGGUA GUAAAAAU | AUUUUUAC CUGAUGA X GAA ACCCAAGA |
| 16 | UGGGUAGUA AAAAUGGU | ACCAUUUU CUGAUGA X GAA ACUACCCA |
| 25 | AAAAUGGUA GCAACCUG | CAGGUUGC CUGAUGA X GAA ACCAUUUU |
| 49 | GGCGAAAUC CUCCAUGU | ACAUGGAG CUGAUGA X GAA AUUUCGCC |
| 52 | GAAAUCCUC CAUGUUCU | AGAACAUG CUGAUGA X GAA AGGAUUUC |
| 58 | CUCCAUGUU CUUUUCCU | AGGAAAAG CUGAUGA X GAA ACAUGGAG |
| 59 | UCCAUGUUC UUUUCCUU | AAGGAAAA CUGAUGA X GAA AACAUGGA |
| 61 | CAUGUUCUU UUCCUUCC | GGAAGGAA CUGAUGA X GAA AGAACAUG |
| 62 | AUGUUCUUU UCCUUCCC | GGGAAGGA CUGAUGA X GAA AAGAACAU |
| 63 | UGUUCUUUU CCUUCCCU | AGGGAAGG CUGAUGA X GAA AAAGAACA |
| 64 | GUUCUUUUC CUUCCCUU | AAGGGAAG CUGAUGA X GAA AAAAGAAC |
| 67 | CUUUUCCUU CCCUUCUU | AAGAAGGG CUGAUGA X GAA AGGAAAAG |
| 68 | UUUUCCUUC CCUUCUUA | UAAGAAGG CUGAUGA X GAA AAGGAAAA |
| | CCUCCCUU CUUAUCCG | CGGAUAAG CUGAUGA X GAA AGGGAAGG |
| 72 | CUUCCCUUC UUAUCCGC | GCGGAUAA CUGAUGA X GAA AAGGGAAG |
| 73 | UCCCUUCUU AUCCGCUG | CAGCGGAU CUGAUGA X GAA AGAAGGGA |
| 75 | CCCUUCUUA UCCGCUGG | CCAGCGGA CUGAUGA X GAA AAGAAGGG |
| 76 | CUUCUUAUC CGCUGGUC | GACCAGCG CUGAUGA X GAA AUAAGAAG |
| 78 | CCGCUGGUC AUUUCAUC | GAUGAAAU CUGAUGA X GAA ACCAGCGG |
| 86 | CCGCOGGOC ACCOCACC | UGGGAUGA CUGAUGA X GAA AUGACCAG |
| 89 | CUGGUCAUU UCAUCCCA | AUGGGAUG CUGAUGA X GAA AAUGACCA |
| 90 | UGGUCAUUU CAUCCCAU | AAUGGGAU CUGAUGA X GAA AAAUGACC |
| 91 | GGUCAUUUC AUCCCAUU | ACUAAUGG CUGAUGA X GAA AUGAAAUG |
| 94 | CAUUUCAUC CCAUUAGU | CGUUAACU CUGAUGA X GAA AUGGGAUG |
| 99 | CAUCCCAUU AGUUAACG | GCGUUAAC CUGAUGA X GAA AAUGGGAU |
| 100 | AUCCCAUUA GUUAACGC | GCGGCGUU CUGAUGA X GAA ACUAAUGG |
| 103 | CCAUUAGUU AACGCCGC | UGCGCGU CUGAUGA X GAA AACUAAUG |
| 104 | CAUUAGUUA ACGCCGCA | GAGGCGAA CUGAUGA X GAA AGCCUUGC |
| 118 | GCAAGGCUA UUCGCCUC | GGGAGGCG CUGAUGA X GAA AUAGCCUU |
| 120 | AAGGCUAUU CGCCUCCC | 000000 |
| 121 | AGGCUAUUC GCCUCCCG | |
| 126 | AUUCGCCUC CCGGGUGU | 1.0.10 |
| 135 | CCGGGUGUU AAAGCCAC | |
| 136 | CGGGUGUUA AAGCCACA | |
| 147 | GCCACAAUC CUCACUAC | |
| 150 | ACAAUCCUC ACUACCCC | 00000 |
| 154 | UCCUCACUA CCCCUCAU | AUGAGGG CUGAUGA X GAA AGUGAGGA |
| 160 | CUACCCUC AUAAUGCC | GGCAUUAU CUGAUGA X GAA AGGGGUAG |
| 163 | CCCCUCAUA AUGCCUUA | UAAGGCAU CUGAUGA X GAA AUGAGGGG |
| 170 | UAAUGCCUU ACUUUUUA | UAAAAAGU CUGAUGA X GAA AGGCAUUA |
| 171 | AAUGCCUUA CUUUUUAG | CUAAAAAG CUGAUGA X GAA AAGGCAUU |
| 174 | GCCUUACUU UUUAGAUC | GAUCUAAA CUGAUGA X GAA AGUAAGGC |
| 175 | CCUUACUUU UUAGAUCU | AGAUCUAA CUGAUGA X GAA AAGUAAGG |
| 176 | CUUACUUUU UAGAUCUA | UAGAUCUA CUGAUGA X GAA AAAGUAAG |
| 177 | UUACUUUUU AGAUCUAC | GUAGAUCU CUGAUGA X GAA AAAAGUAA |
| 178 | UACUUUUUA GAUCUACU | AGUAGAUC CUGAUGA X GAA AAAAAGUA |
| 182 | UUUUAGAUC UACUAUUG | CAAUAGUA CUGAUGA X GAA AUCUAAAA |
| 184 | UUAGAUCUA CUAUUGAC | GUCAAUAG CUGAUGA X GAA AGAUCUAA |
| 187 | GAUCUACUA UUGACGAU | AUCGUCAA CUGAUGA X GAA KGUAGAUC |
| 189 | UCUACUAUU GACGAUGA | UCAUCGUC CUGAUGA X GAA AUAGUAGA |
| 201 | GAUGAUGUU CGAAUUUC | GAAAUUCG CUGAUGA X GAA ACAUCAUC |

Table III

| Nt. Substrate Position 202 AUGAUGUUC GAAUUUCC GGAAAUUC CUGAUGA X GAA AACAUCAU 207 GUUCGAAUUU CCGGAUU AAUCCGGA CUGAUGA X GAA AACAUCAU 208 UUCGAAUUU CCGGAUUU AAAUCCGG CUGAUGA X GAA AAUUCGAA 209 UUCGAAUUUC CGGAUUU AAAUCCGG CUGAUGA X GAA AAUUCGAA 209 UUCGAAUUUC CGGAUUUC GAAUUCC GAAUUCC GAAUUCC GCGAUUC AAAUCGGA CUGAUGA X GAA AAUUCGAA 215 UUCCGGAUUU CCCAUUUC AAAUGGGA CUGAUGA X GAA AAUUCGAA 216 UUCCGAUUUC CCAUUUCC AAAAUGGGA CUGAUGA X GAA AAUCCGGA 217 CCGGAUUUC CCAUUUCCU AAAAUGGGA CUGAUGA X GAA AAUCCGGA 227 UUUCCCAUUU CUUAUCGUU AAGAAUGG CUGAUGA X GAA AAUCCGGA 223 UUUCCCAUUU CUUCUCUU AACAAUGG CUGAUGA X GAA AAUCCGGA 224 UUCCCAUUUC UAUCGUA UUACGAUA CUGAUGA X GAA AAUCGGAA 225 CCAUUUCUU UAUCGUA UACGAUAC CUGAUGA X GAA AAUGGGAA 226 CCAUUUCUU UAUCGUA UACGAUAC CUGAUGA X GAA AAUGGGAA 227 AUUUCAUCGUA CUGUAUCA UACGAUAC CUGAUGA X GAA AAUAGGAA 228 AUUUCAUCGUA CUGUAUCA UACGAUAC CUGAUGA X GAA AAUAGGAA 231 UUCUAUCGUA ACUAUUAA UUAACGAU CUGAACAU AAUAGCACAU 231 UUAACGAU UUAAAUUC CAAUUUAA UUAACGAU CUGAUGA X GAA ACAAUAGA 237 GUAACGAU UUAAAUUC CAAUUUAA UUAACGAU CUGAUGA X GAA ACGAUAGA 238 UUAACUAUU AAUUCCCC GGGAAUU CUGAUGA X GAA ACGAUAGA 237 GUAACUAUU AAUUCCCC GGGAAUU CUGAUGA X GAA ACGAUAGA 238 UUAACUAUU AAUUCCCC GGGAAUU CUGAUGA X GAA ACGAUAGA 238 UUAACUAUA AAUUCCCC GGGAAUU CUGAUGA X GAA ACGAUAGA 241 UAUUAAAUUC CCCCUCUC CACAGGGG CUGAUGA X GAA AUUAAUAAU 241 UAUUAAAUUC CCCCUCUC CACAGGGG CUGAUGA X GAA AUUAAUAA 241 AUUAAAUUC CCCCUCUC CACAGGGG CUGAUGA X GAA AUUUAAUAA 241 AUUAAAUUC CCCCUCUC CACAGGGG CUGAUGA X GAA AUUUAAUAA 241 AUUAAAUUC CCCCUCUC CACAGGGG CUGAUGA X GAA AUUUAAUAA 242 AUAUAAAUUC CCCCUCUC CACAGGGG CUGAUGA X GAA AUUUAAUAA 243 AUUAAAUUC CCCCCUCUC CACAGGGG CUGAUGA X GAA AUUUAAUAA 244 AUUACAAAUAU CCCCCUCUC CACAGGGG CUGAUGA X GAA AUUUAACAA 245 AGAUGACAAUAA CUGAGAA CUGAGA | | | |
|--|-----|--------------------|---------------------------------|
| POSITION 202 AUGAUGUU GAAUUUCC GGAAUUU CUGAUGA X GAA AACAUCAU 208 UUCGAAUUU CCCGGAUUU AAUCCGGA CUGAUGA X GAA AUUCGAAC 208 UUCGGAUUU CCGGAUUU AAAUCCGGA CUGAUGA X GAA AUUCGAAC 209 UCCGAUUUU CGGAUUUC GAAAUCCG CUGAUGA X GAA AAUUCGAAC 215 UUCCGGAUUU CCCAUUU AAAUGGGA CUGAUGA X GAA AAUCCGGA 215 UUCCGGAUUU CCCAUUUC GAAAUGGG CUGAUGA X GAA AAUCCGGA 217 CCGGAUUUC CCAUUUC GAAAUGGG CUGAUGA X GAA AAUCCGGA 217 CCGGAUUUC CCAUUUC GAAAUGG CUGAUGA X GAA AAUCCGGA 217 CCGGAUUUC CUAUCGUA GAAAUGG CUGAUGA X GAA AAUCCGGA 222 UUUCCCAUUU CUAUCGUA UCACAUCCU GAAAUGG CUGAUGA X GAA AAUCCGGA 223 UUCCCAUUUC UAUCGUA UACGAUGA CUGAUGA X GAA AAUCCGGA 224 UCCCAUUUC DAUCGUAA UACGAUGA CUGAUGA X GAA AAUCGGAA 224 UCCCAUUUC DAUCGUAA CUGAUGA CUGAUGA X GAA AAUCGGAA 225 CCAUUUCUD DACGUAACUA DACGAUGA CUGAUGA X GAA AAUGGGAA 226 CCAUUUCUD DACGUAACUA DAGGUACA CUGAUGA X GAA AAUGGGAA 227 UCUACCGAUU CUAUCGUA GUAACUAU CUGAUGA X GAA AAUGGGAA 228 AUUUCAUC GUAACUAU AUGAGUAC CUGAUGA X GAA AAUGGGAA 238 UCAACUAU AAUUUCA CUGAACUAU CUGAUGA X GAA AAUGGGAA 237 GUAACUAUU AAUUUCCC GGGAAUUU CUGAUGA X GAA AAUGGGAA 238 UAACUAUUA AAUUUCCC GGGAAUUU CUGAUGA X GAA AAUGUUAC 238 UAACUAUUA AAUUUCCCC GGGAAUUU CUGAUGA X GAA AAUGUUAC 238 UAACUAUUA AAUUUCCCC GGGGAAUU CUGAUGA X GAA AAUGUUAC 2412 UAUUAAAUUC CCCUCUGC GAGGGG CUGAUGA X GAA AAUGUUAC 2412 UAUUAAAUUC CCCUCUGC GAGGGG CUGAUGA X GAA AAUGUUACA 2413 AUUAAAUUC CCCUCUGC GAGGGG CUGAUGA X GAA AAUGUUAC 2414 UAUUAAAUU CCCCUCUGC GAGGGG CUGAUGA X GAA AAUGUUAC 2415 AAUUUAAUU CCCCUCUGC GAGGGG CUGAUGA X GAA AAUGUUAC 2416 AUUCCCCU UCCUGCC GAGGGG CUGAUGA X GAA AAUGUUAC 2417 AUUAAAUU CCCCCCUCUG CAGGGGG CUGAUGA X GAA AAUGUUAC 2418 AUUCCCCU UCCUCUC GAGGGG CUGAUGA X GAA AAUUUAUU 2418 AUUCCCCU UCCUCUC GAGGGG CUGAUGA X GAA AAUGUUAC 2419 CUUUAACUU CCCCCCCG CAGAGGG CUGAUGA X GAA AAUGUUAC 2410 AAUUAAAUU CCCCCCCGC CAGAGGG CUGAUGA X GAA AAUUUAUAU 2411 AAUAAAUU CCCCCCCGGAAC CUCAGAC CUGAUGA X GAA AAUUUAUAU 2411 AAUAAAUU GCCCCCC GAGAGGC CUGAUGA X GAA AAUUUAAU 2411 AAUAAAUU GCCCCCCG CAGAGGG CUGAUGA X GAA AAUUUAAU 2412 AAUACUAC AAUACAC CUCAGAC CUGAUGA X GAA AAUUU | Nt. | Substrate | Ribozyme |
| 202 AUGAGGUU GAAUUU CCGGAUU AAUCCGG CUGAUGA X GAA AGUUGGAA CO COGAUGA X GAA AGUUGGAA CO COGAUGU UCCGAUUU AAUCCGG CUGAUGA X GAA AGUUGGAA CO COGAUGU UCCGAUUU AAUCCG CUGAUGA X GAA AGUUGGAA CO COGAUGU UCCGAUUU AAUCCG CUGAUGA X GAA AAUUCGAA CO COGAUGU UCCGAUUU AAAUGGG CUGAUGA X GAA AAUUCGGA COGAUUU COCAUUU AAAUGGG CUGAUGA X GAA AAUCCGG AC CUGAUGA X GAA AGUCGGAA COGAUGA X GAA AGUGGGAA COCAUUU CUUAUGGU ACGAUGA X GAA AGUGGGAA COGAUGA X GAA AGUGGGAA COCAUUUCU ACCAUUUCU ACCAUUCA ACGAUGA X GAA AGUGGGAA COCAUUCAA COGAUGA X GAA AGUGGGAA COCAUUCAA COCAUCAA X GAA AGAAGGGAA COCAUCAA COCAUCAA COCAUCAA COCAUCAA X GAA ACAAGAGGA COCAUCAA COCAUCAA COCAUCAA COCAUCAA COCAUCAA X GAA ACAAGAGAA COCAUCAA COCAUCAA X GAA ACAACAGAA COCAUCAA COCAUCAA COCAUCAA X GAA ACAACAGAA COCAUCAA COCAUCAA X GAA ACAACAGAA COCAUCAA COCAUCAA X GAA ACAACAGAA COCAUCAA COCAUCAA X GAA ACAACACACACACACACACACACACACACACACAC | | - | • |
| DOTO DUCGAAUUU CCGGAUUU DUCGAAUUUC CGGAUUUC GGAUUUC GAAUCGGC CUGAUGA X GAA AAUUCGGA 215 UUCCGGAUU CCCAUUUC AGAAUGGG CUGAUGA X GAA AAUCCGGA 217 CCGGAUUUC CAUUUCU AGAAUGGG CUGAUGA X GAA AAUCCGGA 217 CCGGAUUUC CAUUUCU AGAAUGGG CUGAUGA X GAA AAUCCGGA 222 UUUCCCAUUUC AGAAUGGG CUGAUGA X GAA AAUCCGGA 223 UUUCCCAUUUC GAGAUGG CUGAUGA X GAA AAUCCGGA 224 UUCCCAUUUC UAUCGUA 225 UUUCCCAUUU UAUCGUA 226 CCAUUUCUA UCGUAACU AUCCGAUGA CUGAUGA X GAA AAUGGGAA 227 UCCCAUUUCU UAUCGUAA 228 AUUUCUAUC UAUCGUAA 229 AUUUCUAUC UAUCGUAA 229 AUUUCUAUC UAUCGUAA 220 AUUUCUAUC UAUCGUAA 221 UCUAUCGUA UCGUAACU AGUUACGA CUGAUGA X GAA AAUGGGAA 221 UCUAUCGGUA ACUAUUAA 222 AUUUCUAUC UAAAUUC CGAAUUUCAAUC GAAUUUAA 223 UCUAUCGGUA ACUAUUAA 223 UCUAUCGGUA ACUAUUAA 224 UUAAAUUC CGAAUUUAAAUUC CGAGAUUU CAUCAGAA X GAA ACGAUAGA 223 UAACUAUUA AUUAAAUUC CGAGAAUUU COAUGA X GAA ACGAUAGA 224 UAUAAAUUC CCCUCUGC CGGGAAUUU COAUGAA X GAA ACUAUAAA 224 UAUAAAUUC CCCUCUGC CACAGGGG CUGAUGA X GAA ACUAUAAA 224 UAAAAUUU UAACUCC CCCUCUGC CACAGGGG CUGAUGA X GAA ACUCUCA 225 CUGAGGGA CUGAUGA X GAA ACUCUCA 226 CUGAGGGA CUGAUGA X GAA ACUCUCA 227 CACAGGGA CUGAUGA X GAA ACUCUCA 228 CUGAGGGA CUGAGGA X GAA ACUCUCA 228 CUCAAGGC CUGAUGA X GAA ACUCUCA 229 CUUUAACUC CCCUCUCA ACUAUAAAAAUU ACUCCCUCUCA CACAGGGG CUGAUGA X GAA ACUCUCA 229 CUUUAACUC UACCAGAGGU CUGAUGA X GAA ACUCUCA 229 CUUUAACUC UACCAGAGGC CUGAUGA X GAA ACUCUCA 229 CUUUAACUC UACCAGAGG CUGAUGA X GAA AAGUCUAA 229 CUUUAACUC UACCAGAGG CUGAUGA X GAA AAGUCUAA 230 CACACCUUA ACUCUCU AAC | | AUGAUGUUC GAAUUUCC | GGAAAUUC CUGAUGA X GAA AACAUCAU |
| DOB UUCGAAUUU CCGAUUUC AAAUCCG CUGAUGA X GAA AAUCCGAU 209 UCGAAUUUC CGAUUUC GAAAUCCG CUGAUGA X GAA AAUCCGGA 216 UUCCGGAUU CCCAUUUC GAAAUCCG CUGAUGA X GAA AAUCCGGA 216 UCCGGAUUC CCAUUUCU GAAAUGGG CUGAUGA X GAA AAUCCGGA 217 CCGGAUUUC CCAUUUCU ACGAUGA CUGAUGA X GAA AAUCCGGA 218 CUCCCAUUU CUAUCGU ACGAUGA CUGAUGA X GAA AAUCCGG 222 UUUCCCAUU CUAUCGU ACGAUGA CUGAUGA X GAA AAUCGG 223 UUCCCAUUU CUAUCGU ACGAUGA CUGAUGA X GAA AAUCGGG 224 UUCCCAUUUC UAUCGUA ACGAUGA CUGAUGA X GAA AAUGGGAA 224 UUCCCAUUUC UAUCGUA ACGAUGA CUGAUGA X GAA AAUGGGAA 225 CUCCUUUC UUCGUAACU AUCGUAA AUCAGAUA CUGAUGA X GAA AAUGGGAA 226 CCCAUUUCUA UCGUAACU AUCAGAUA CUGAUGA X GAA AAUGGGAA 227 AUUUCAAUCGUA AUCAGAUA CUGAUGA X GAA AAUGGGAA 228 AUUUCUAUC GUAACUA UUAAUAGU CUGAUGA X GAA AUGAAAU 231 UUCAGAUGA WAAUUCCC GGGAAUUU CUGAUGA X GAA ACGAUGA 231 UUCAGAUGA WAAUUCCC GGGAAUUU CUGAUGA X GAA AUGAAAU 233 UUAACUAUUA AAUUCCC GGGAAUUU CUGAUGA X GAA AUGAGAU 233 UUAACUAUUA AAUUCCC GGGAAUUU CUGAUGA X GAA AUGAGAU 234 UUAUAAAUUC CCCUCUGG CAGAGGG CUGAUGA X GAA AUGAGAU 243 AUUAAAUUC CCCUCUGG CAGAGGGG CUGAUGA X GAA AUGUAAA 244 UUCAGAGAGCUU AACCUCUG CAGAGGGG CUGAUGA X GAA AUGUAAA 258 GCUGAAGUU GAGAGCUU AACCUCU CAGAGAU CUGAUGA X GAA AUGUAACA 258 GCUGAAGUU GAGAGCUU AACCUCU CAGAGGGA CUGAUGA X GAA AUGUCAGC 263 ACGUGGGU GCCCGAAAG CUUCAGCA CUGAUGA X GAA ACGUCUCA 264 UUGAGAGCUU AACCUCU AACCUCU AACCUCU CAGAGUU CUGAUGA X GAA ACCUCACACU 265 GAGAGCUUU AACCUCU CAGAGGU CUGAUGA X GAA AACCUCUC 266 AAGGCUUUA ACCUCU AACCUCU AACCUCU CAGAGUU CUGAUGA X GAA AACCUCUC 267 GAAGGAUU CUGAGCA CUGAUGA X GAA AACCUCUC 268 AACGUUUAA CUCUCUCU AACCUCU AACCUCUAAA CUCAUGA X GAA AACCUCUC 269 CUGUAACA CUCAUGA X GAA AACCUCUC AACCUCUAAA CUCAUGA X GAA AACCUCUCA 279 UUCACACUU CUCUCUA AACCUCU CACAGAG CUCA | | | AAUCCGGA CUGAUGA X GAA AUUCGAAC |
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| 237 GUACUAUU AAGUUCCC GGGAAUU CUGAUGA X GAA AUGUUAC 238 UAACUAUUA AABUUCCCC GGGAAUU CUGAUGA X GAA AAUAGUUA 242 UAGUAAAAUU CCCCUCUG CAGAGGG CUGAUGA X GAA AAUAGUUA 243 AUUAAAUUC CCCUCUGC GCAGAGGG CUGAUGA X GAA AAUUUAAU 248 AUUCCCCCU UGCUGAAG CUUCAGCA CUGAUGA X GAA AGUUCAGC 258 GCUGAAGUU GGGUUGCC GGCAACCC CUGAUGA X GAA ACUUCAGC 258 GCUGAAGUU GGGUUGCC GGCAACCC CUGAUGA X GAA ACUUCAGC 263 AGUUGGGUU GCCUGAAG CUUCAGGC CUGAUGA X GAA ACUUCAGC 276 GAAGGAAUU GAGAGCUU AAGCCUC CUGAUGA X GAA ACUUCAGC 284 UGAGAGCUU AACUCUGC CAGAGUUA CUGAUGA X GAA ACUCCUC 285 GAGAGCUUU AACUCUGC GCAGAGUU CUGAUGA X GAA AGCUCUCA 285 GAGAGCUUU AACUCUGC GCAGAGU CUGAUGA X GAA AGCUCUCA 286 AAGCUUUA ACUCUGCC GCAGAGU CUGAUGA X GAA AAGCUCUC 290 CUUUAACU CUCCCCACUU AAGUGCA CUGAUGA X GAA AAGCUCUC 290 CUUUAACU CACCUGAAA UUCAGGU CUGAUGA X GAA AAGCUCUC 290 UGCCACUU CACCUGAAA UUCAGGU CUGAUGA X GAA AGGUCUC 291 UGCCACUU CACCUGAAA UUCAGGU CUGAUGA X GAA AGGCAUU 313 AAAGGCCUC AUAAAAUU AAUUUUU AAAAAUUUU CUGAUGA X GAA AGGCAUU 316 UGCCUCAUA AAAUUUUU AAAAAUUU CUGAUGA X GAA AGGCAUU 321 CAUAAAAUU UUUAUGCC GCAGAGAA CUGAUGA X GAA AGGCAUU 322 AUAAAAUU UUUAUGCC GAGCAUAA CUGAUGA X GAA AGGCAUU 322 AUAAAAUU UUUAUGCC GAGCAUAA CUGAUGA X GAA AUUUUAU 323 UAAAAAUUU UUAUGCCU AGCAAAA CUGAUGA X GAA AUUUUAU 324 AAAAUUUUU UUAUGCUC GAGCAUAA CUGAUGA X GAA AAUUUUAU 325 AUAAAAUUU UUAUGCUC AAGCAUAA CUGAUGA X GAA AAUUUUAU 326 AAAAUUUUU AAAAAUU AAAAUUUU AAAAAUUU AAAAAUUUAAAA CUGAUGA X GAA AAAUUUAAA 323 UAAAAAUUU UUAUGCUC AAGCAAAA CUGAUGA X GAA AAAUUUAAA 324 AAAAUUUUU AAAAAC CUGAUGA X GAA AAAAUUUAAAA 335 UAUGCCUCUU AAGAGAA CUGAUGA X GAA AAAAUUUAA 336 AAAAUUUUU AAAAAAC CUGAUGA X GAA AAAAUUUAA 337 CUCUUUCUC UACAAAA CUGAUGA X GAA AAAAAUUAAA 337 CUCUUCUCU CUCUAAAA UUGAGAA CUGAUGA X GAA AAAAAUUAAA 338 UAUGCCUCU CUCAAAAA CUGAUGA X GAA AAAAAAAAAAAAAAAAAAAAAAAAAAA | | UCGUAACUA UUAAAUUC | |
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| 242 UAUUAAAUU CCCUCUGC CAGAGGGG CUGAUGA X GAA AUUUUAAU 243 AUUAAAUU CCCUCUGC GCAGAGGG CUGAUGA X GAA AAUUUUAAU 248 AUUCCCCUC UGCUGAAG CUUCAGCA CUGAUGA X GAA AAUUUUAAU 258 GCUGAAGUU GGGUGCC GGCAACCC CUGAUGA X GAA ACUCCAGC 263 AGUUGGGUU GCCUGAAG CUUCAGGC CUGAUGA X GAA ACUCCAGC 264 GCAGAGAUU GAGAGCUU AAGCUCUC CUGAUGA X GAA ACUCCACACU 276 GCAGAGAUU AACUCUGC CAGAGUUA CUGAUGA X GAA ACUCCUCA 284 UGAGAGCUU AACUCUGC CAGAGUUA CUGAUGA X GAA ACUCCUCA 285 GACAGCUUU AACUCUGC GCACAGUU CUGAUGA X GAA AGCUCUCA 286 AGAGCUUU AACUCUGC GCACAGUU CUGAUGA X GAA AGCUCUCA 286 AGAGCUUU AACUCUGC GCACAGGU CUGAUGA X GAA AGCUCUCA 290 CUUUAACUC UGCCACUU AAGUGGCA CUGAUGA X GAA AGCUCUCA 298 CUGCCACUU CACCUGAAA UUCAGGUG CUGAUGA X GAA AGUUAAAG 299 UGCCACUU CACCUGAAA UUCAGGU CUGAUGA X GAA AGUUAAAG 299 UGCCACUU CACCUGAAA UUCAGGU CUGAUGA X GAA AGUGAGCA 299 UGCCACUU AAAAAUU AAUUUUU CUGAUGA X GAA AGGCAUU 313 AAAUGCCUC AUAAAAUU AAUUUUUU CUGAUGA X GAA AGGCAUU 316 UGCCUCAUA AAAUUUUU AAAAAUUU CUGAUGA X GAA AGGCAUU 321 CAUAAAAUU UUUAUGC GCAGAGAA CUGAUGA X GAA AGGCAUU 322 AUAAAAUUU UUUAUGC GCAGAGAA CUGAUGA X GAA AAUUUUAU 322 AUAAAAUUU UUAUGCC GAGCAUAA CUGAUGA X GAA AAUUUUAU 323 UAAAAUUUU UUAUGCC GAGCAUAA CUGAUGA X GAA AAUUUUAU 324 AAAAUUUUU UAUGCUC AGCAAAA CUGAUGA X GAA AAUUUUAU 325 AAAUUUUUU UAUGCUC AAGACAUA CUGAUGA X GAA AAAUUUUAU 326 AAUUUUUU UAUGCUC AAGACAUA CUGAUGA X GAA AAAUUUUA 327 AAAAUUUUU UAUGCUC AAGAGAA CUGAUGA X GAA AAAUUUUA 328 AAAAUUUUU UAUGCUC AAGAGAA CUGAUGA X GAA AAAAUUUAA 331 UUUAUGCUC UUCUCUCU AAGAGAA CUGAUGA X GAA AAAAUUUAA 332 WAAAAUUUU UAUGCUCU AAGAGAA CUGAUGA X GAA AAAAUUUAA 333 UAUGCCUCUU AAGAGAA CUGAUGA X GAA AAAAUUUAA 334 AUGCUCUU UCUCUCUA AAGAGCA CUGAUGA X GAA AAAAAUU 344 AAAAUUUUU UCUACAA UUUGAAAA CUGAUGA X GAA AAAAAUU 355 AAAUUUUUA UCUCUCUA CAAAAGA CUGAUGA X GAA AAAAAUU 366 AAUUUUUCU UCUCUCUA CAAAAGA CUGAUGA X GAA AAAAAUU 377 CUCUUUCUC UUCUACAA UUUCUAAAA CUGAUGA X GAA AAAAAUU 378 CUCUUCUCU UACAAAAA CUGAUGA X GAA AAAAAAAAAAAAAAAAAAAAAAAAAAA | | UAACUAUUA AAUUCCCC | 0000 |
| 243 AUUAAAUUC CCCUCUGC GCAGAGGG CUGAUGA X GAA AAUUCAAU 248 AUUCCCCUC UGCUGAAG 248 AUUCCCCUC UGCUGAAG 258 GCUGAAGUU GGGUUGCC GCAACC CUGAUGA X GAA ACUUCAGC 263 AGUUGGGUU GCCUGAAG 276 GAAGGAAUU GAGACCUU AAGCUUC CUGAUGA X GAA ACCCAACU 276 GAAGGAAUU GAGACCUU CAGAGG CUUCAGGC CUGAUGA X GAA ACCCAACU 276 GAAGGAUU AACUCUG CAGAGUU CUGAUGA X GAA AUUCCUUC 284 UGAAGGCUU AACUCUGC GCAGAGUU CUGAUGA X GAA AGCUCUC 285 GAGACCUUU AACUCUGC GCAGAGUU CUGAUGA X GAA AAGCUCUC 286 AGAGCUUUA ACUCUGC GCAGAGUU CUGAUGA X GAA AAGCUCUC 286 AGAGCUUUA ACUCUGC GCAGAGUU CUGAUGA X GAA AAGCUCUC 290 CUUUAACUC UGCCACUU AAGUGGCA CUGAUGA X GAA AAGUCUC 290 UGCCACUU CACCUGAAA UUUCAGGU CUGAUGA X GAA AAGUCACA 299 UGCCACUU CACCUGAA UUUCAGGU CUGAUGA X GAA AAGUGGCA 299 UGCCACUU CACCUGAA UUUCAGGU CUGAUGA X GAA AAGUGGCA 311 AAAGGCCUC AUAAAAUU AAUUUUAU CUGAUGA X GAA AAGUGAGA 312 CAUAAAAUU UUUAUGCC GCAGAGAU CUGAUGA X GAA AAGUGGCA 321 CAUAAAAUU UUUAUGCC GCAGAAA CUGAUGA X GAA AAUUUUAU 322 AUAAAAUUU UUUAUGCC GCAGAAAA CUGAUGA X GAA AAUUUUAU 322 AUAAAAUUU UUUAUGCC GAGCAUAA CUGAUGA X GAA AAUUUUAU 323 UAAAAUUU UUAUGCUC AGCAUAAA CUGAUGA X GAA AAUUUUAU 324 AAAAUUUUU AUGCUCU AGAGCAUA CUGAUGA X GAA AAAUUUUA 325 AAAUUUUUU AUGCUCU AAGACAU CUGAUGA X GAA AAAUUUUA 326 AAUUUUUU AUGCUCU AAGACAU CUGAUGA X GAA AAAUUUUA 327 AAAAAUUU UUUAUGCUCU AAGACAU CUGAUGA X GAA AAAAUUUU 328 AAAAUUUUU AUGCUCUU AAGAGCAU CUGAUGA X GAA AAAAUUUU 329 AAAAAAUUU AUGCUCUU AAGAGACA CUGAUGA X GAA AAAAAUUU 320 AAAAAUUUU AUGCUCUU AAGAGAA CUGAUGA X GAA AAAAAUUU 331 UUUAUGCUCU UUCUCUU AAGAGAA CUGAUGA X GAA AAAAAUUU 333 UAUGCUCUU CUCUUCUA AAGAGCA CUGAUGA X GAA AAAAAUUU 333 UAUGCUCUU CUCUUCUA AAGAGCA CUGAUGA X GAA AAAAAUUU 333 UAUGCUCUU CUCUUCUA AAGAGCA CUGAUGA X GAA AAAAAUUU 333 UAUGCUCUU CUCUUCAAA CUGAUGA X GAA AAAAAUUU 334 AUGCCUCUU CUCAAAAA CUGAUGA X GAA AAAAAUUU 335 OGCUCUUUC UACAAAAA CUGAUGA X GAA AAAAAUUU 336 AAGCAAAA CUGAUGA X GAA AAAAAUUU 337 OCCUUUCUA CAAAAAG CUGAUGA X GAA AAAAAUUU 340 UUUCAGAAA CUGAUGA X GAA AAAAAUUU 340 UUUCACAAAA CUGAUGA X GAA AAAAAUUUAACAA CUGAUGA X GAA AAAAAUUUAAAAAAUUU CUGAAAAA CUGA | | UAUUAAAUU CCCCUCUG | 0.10.10000 |
| 248 AUUCCCCUC UGCUGAAG CUUCAGCA CUGAUGA X GAA AGGGGAAD 258 GCUGAAGUU GGCUGCC GGCAACCC CUGAUGA X GAA ACUCCAGC 263 AGUUGGGUU GCCUGAAG CUUCAGGC CUGAUGA X GAA ACCCAACU 276 GAAGGAAUU GAGAGCUU AAGCUCUC CUGAUGA X GAA ACCCAACU 276 GAAGGAAUU GAGAGCUU AAGCUCUC CUGAUGA X GAA AGCUCUCA 284 UGAGAGCUU AACUCUGC CAGAGUU CUGAUGA X GAA AGCUCUCA 285 GAGAGCUUU ACUCUGC CAGAGUU CUGAUGA X GAA AAGCUCUC 286 AGAGCUUU ACUCUGCC GGCAGAGUU CUGAUGA X GAA AAGCUCUC 287 GAGAGCUUU ACUCUGCC GGCAGAGUU CUGAUGA X GAA AAGCUCUC 288 AGAGCUUUA ACUCUGCC GGCAGAGUU CUGAUGA X GAA AAGCUCUC 2890 CUUUAACUC UGCCACUU AAGUGGCGC CUGAUGA X GAA AGGUAAC 2990 UGCCACUUC ACCUGAA UUUCAGGUG CUGAUGA X GAA AGUGACA 2990 UGCCACUUC ACCUGAA UUUCAGGUG CUGAUGA X GAA AGUGACA 2990 UGCCACUUC ACCUGAAA UUUCAGGUG CUGAUGA X GAA AGUGACA 313 AAAUGCCUC AUAAAAUU AAUUUUUU AAGAGUC CUGAUGA X GAA AGUGAGCA 316 UGCCACUU ACCUGAAA UUUUAAGGU CUGAUGA X GAA AAGUGGCA 321 CAUAAAAUU UUUUAUGC CCAUAAAA CUGAUGA X GAA AUUUUAUA 322 AUAAAAUUU UUUAUGCU AGCACUAAA CUGAUGA X GAA AAUUUUAA 323 UUAAAAUUUU UUUAUGCU AGGACAUA CUGAUGA X GAA AAAUUUUA 324 AAAAUUUUU AUGCCUCU AAGAGAAA CUGAUGA X GAA AAAUUUUA 325 AAAUUUUUU AUGCUCUU AAGAGCAU CUGAUGA X GAA AAAUUUUA 326 AAAUUUUUU AUGCUCUU AAGAGCAU CUGAUGA X GAA AAAUUUUA 327 AAAAUUUUU AUGCUCUU AAGAGCAU CUGAUGA X GAA AAAUUUUA 328 AAAUUUUUU AUGCUCUUU AAGAGCAU CUGAUGA X GAA AAAAUUUU 329 AAAAUUUUU AUGCUCUUU AAGAGCAU CUGAUGA X GAA AAAAUUUU 320 AAAAUUUUU AUGCUCUUU AAGAGAAA CUGAUGA X GAA AAAAUUUUA 331 UUUAUGCU UUCUUCUU AAGAGAAA CUGAUGA X GAA AAAAUUUUA 332 UUGAUGAU CUCUUCUU AAGAGAAA CUGAUGA X GAA AAAAUUUUA 333 UAUGCCUCUU CUCUUCUU AAGAGAAA CUGAUGA X GAA AAAAUUUAA 334 AUGCUCUUU UCUCUUCU AAGAGAAA CUGAUGA X GAA AAAAAUU 335 UGCUUUUCU UUCUACAA UUGUAGAA CUGAUGA X GAA AAAAAUU 336 OGUUUCUCUU UACAAAAA 337 CUCUUUCUU UACAAAAA 338 OGUUCACCAA UUGUAGAA CUGAUGA X GAA AAAAAUU 339 CUUUCUCUU CUACAAAAA CUGAUGA X GAA AAAAAAAAAAAAAAAAAAAAAAAAAAA | | AUUAAAUUC CCCUCUGC | GONONICCE COCKETO |
| 258 GCUGARGUU GGGUUGCC GGCAACCC CUGAUGA X GAA ACUCAGCC 263 AGUUGGGUU GCCUGAAG CUUCAGGC CUGAUGA X GAA ACCCAACU 276 GAAGGAAUU GAGAGCUU AAGCUCUC CUGAUGA X GAA ACCUCUC 284 UGAGAGCUU NACUCUG CAGAGUU CUGAUGA X GAA AGCUCUC 285 GAGAGCUUU AACUCUGC GCAGAGUU CUGAUGA X GAA AAGCUCUC 286 AGAGGCUUU ACUCUGCC GCAGAGUU CUGAUGA X GAA AAGCUCUC 290 CUUUAACUC UGCCCACUU AAGUGGCA CUGAUGA X GAA AAGCUCUC 299 UGCCACUU CACCUGAA UUCAGGUG CUGAUGA X GAA AAGCUCU 299 UGCCACUU CACCUGAA UUCAGGUG CUGAUGA X GAA AAGCUCU 313 AAAUGCCUC AUAAAAUU AAUUUUUAU CUGAUGA X GAA AAGCAUU 316 UGCCUCAUA AAAUUUUU AAAAAUUU CUGAUGA X GAA AAGCAUAA 317 CAUAAAAUU UUUAUGCU GCAGAGAU CUGAUGA X GAA AUUUUAAG 322 AUAAAAUUU UUUAUGCU GAGCAUAAA CUGAUGA X GAA AUUUUAA 323 UAAAAUUUU UUAUGCU GAGCAUAAA CUGAUGA X GAA AAUUUUAA 324 AAAAUUUUU UAUGCUCU AGAGCAUA CUGAUGA X GAA AAAUUUUAA 325 AAAUUUUUU UAUGCUCU AGAGCAUA CUGAUGA X GAA AAAUUUUAA 326 AAAUUUUUU UAUGCUCU AAGAGCAU CUGAUGA X GAA AAAAUUUUA 327 AAAAUUUUU UAUGCUCU AAGAGCAU CUGAUGA X GAA AAAAUUUAA 328 AAAUUUUUU AUGCUCUU AAGAGCAU CUGAUGA X GAA AAAAUUUUA 329 AAAAUUUUUU AUGCUCUU AAGAGCAU CUGAUGA X GAA AAAAAUUUA 320 AAAAUUUUUU AUGCUCUU AAGAGCAU CUGAUGA X GAA AAAAAUUUA 321 AAAAUUUUU UAUGCUCU AAAAAAA CUGAUGA X GAA AAAAAUUUA 322 AAAAUUUUUU AUGCUCUU AAGAGCAU CUGAUGA X GAA AAAAAUUUA 323 UAUGCUCUU CUUCUUU AAGAGCA CUGAUGA X GAA AAAAAUUU 324 AAAAUUUUU UAUGCUCUU AAGAGCAU CUGAUGA X GAA AAAAAUUU 325 AAAUUUUUU AUGCUCUU AAGAGAGA CUGAUGA X GAA AAAAAUUU 333 UAUGCUCUU CUUUCUAA AAAAGCA CUGAUGA X GAA AAAAAUUU 333 UAUGCUCUU CUUUCUAA AAAAGA CUGAUGA X GAA AAAAAUUU 333 UAUGCUCUU CUUUCUAA UAGAAGAA CUGAUGA X GAA AAAAAUUU 333 UAUGCUCUU CUUCUAA AAAAGA CUGAUGA X GAA AAAAAAUAAAAAAAAAAAAAAAAAAAA | | AUUCCCCUC UGCUGAAG | |
| 263 AGUUGGGUU GCCUGAAG CUUCAGGC CUGAUGA X GAA ACCAACU 276 GAAGGAAUU GAGAGCUU AAGCUCUC CUGAUGA X GAA AUCCAUCA 284 UGAGAGCUU WACCUCUGC CAGAGUUA CUGAUGA X GAA AGCUCUC 285 GAGAGCUUW AACUCUGC GCAGAGUU CUGAUGA X GAA AAGCUCUC 286 AGAGCUUWA ACUCUGCC GCCAGAGUU CUGAUGA X GAA AAGCUCUC 290 CUUWAACUC UGCCACUU AAGUGCA CUGAUGA X GAA AAGCUCUC 290 CUUWAACUC UGCCACUU AAGUGCA CUGAUGA X GAA AGUUAAAG 298 CUGCCACUU CACCUGAA WACAGGGCA CUGAUGA X GAA AGUUAAAG 299 UGCCACUUC ACCUGAAA WACAGGGCA CUGAUGA X GAA AGUGCAG 313 AAAUGCCUC AWAAAAWU AAAAAUU CUGAUGA X GAA AGUGCAG 316 UGCCUCAWA AAAWWA AAAAAWU CUGAUGA X GAA AGUGGCAG 321 CAWAAAAWU WUWAUGC GCAWAAAA CUGAUGA X GAA AGGAGWA 322 AWAAAAWU WUWAUGCU GAGCAWAA CUGAUGA X GAA AAWWUWA 323 WAAAAWWW WUWAUGCUC GAGCAWA CUGAUGA X GAA AAWWWWA 324 AAAAWWWW WUAUGCUC GAGCAWA CUGAUGA X GAA AAWWWWA 325 AAAWWWWW WAUGCUCU AAGAGCAW CUGAUGA X GAA AAAWWWW 326 AAAWWWWW WAUGCUCU AAGAGCAW CUGAUGA X GAA AAAWWWW 327 AAAWWWWW WAUGCUCU AAGAGCAW CUGAUGA X GAA AAAWWWW 328 AAAWWWWW WAUGCUCU AAGAGCAW CUGAUGA X GAA AAAAWWWW 331 WAUGCUCUW WAGCUCUW AAGAGCAW CUGAUGA X GAA AAAAWWWW 333 WAUGCUCUW WAGCUCUW AAGAGCA CUGAUGA X GAA AAAAWWWW 333 WAUGCUCUW CUCUUCUW AAGAGCA CUGAUGA X GAA AAAAAWWWW 333 WAUGCUCUW CUCUUCUW AAGAGCA CUGAUGA X GAA AAAAAWWWWWWAAAAAA CUGAUGA X GAA AAAAAWWWWWAAAAAAAAWWWWWAAAAAAAWWWWAAAAAA | | GCUGAAGUU GGGUUGCC | |
| 276 GAAGGARUU GAGAGCUU AAGCUCUC CUGAUGA X GAA AGCUCUCA 284 UGAAGAGCUU AACUCUG CAGAGUUA CUGAUGA X GAA AGCUCUCA 285 GAGACUUU AACUCUGC GCAGAGUU CUGAUGA X GAA AAGCUCUC 286 AGAGCUUU ACUGCC GCAGAGUU CUGAUGA X GAA AAGCUCUC 290 CUUUAACUC UGCCACUU AAGUGGCA CUGAUGA X GAA AAGCUCUC 290 CUUUAACUC UGCCACUU AAGUGGCA CUGAUGA X GAA AAGUGAAAG 298 CUGCCACUU CACCUGAA UUCAGGUG CUGAUGA X GAA AGUGGCAG 299 UGCCACUUC ACCUGAA UUUCAGGU CUGAUGA X GAA AAGUGGCAG 313 AAAUGCCUC AUAAAAUU AAUUUUAU CUGAUGA X GAA AAGUGGCAG 314 CUGACCAUA AAAUUUUU AAAAAUU CUGAUGA X GAA AGGCAUUU 316 UGCCACUUC ACCUGAAA UUUCAGGU CUGAUGA X GAA AAGUGGCA 321 CAUAAAAUU UUUUAUGC GCAUAAAA CUGAUGA X GAA AUUUUAUG 322 AUAAAAAUUU UUUAUGCU ACCAUAAA CUGAUGA X GAA AUUUUUAU 323 UAAAAAUUUU UUUAUGCU ACCAUAAA CUGAUGA X GAA AAUUUUAU 324 AAAAUUUUU AUUGCUC GAGCAUAA CUGAUGA X GAA AAAUUUUAU 325 AAAUUUUUU AUGCUCU ACAGACAU CUGAUGA X GAA AAAAUUUU 326 AAAUUUUUU AUGCUCU AACAGCAU CUGAUGA X GAA AAAAUUUU 327 AAAAUUUUU AUGCUCUU AACAGACAU CUGAUGA X GAA AAAAUUUU 328 AAAAUUUUU AUGCUCUU AACAGACAU CUGAUGA X GAA AAAAAUUU 339 UUUAUGCUC UUUCUCUU AACAGACAU CUGAUGA X GAA AAAAAUUU 331 UUUAUGCUC UUUCUCUU AACAGACAU CUGAUGA X GAA AAAAAAUUU 333 UAUGCUCUUU CUCUUCUU AACAGACA CUGAUGA X GAA AAAAAAUUU 333 UAUGCUCUU UCUCUUCUU AACAGACAU CUGAUGA X GAA AAAAAAAUU 333 UAUGCUCUU UCUCUUCUA CAAAAGAC CUGAUGA X GAA AAAAAAAAAAAAAAAAAAAAAAAAAAA | | AGUUGGGUU GCCUGAAG | |
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| 326 AAUUUUUUA UGCUCUUU AAAGAGCA CUGAUGA X GAA AAAAAAUU 331 UUUAUGCUC UUUCUCUU AAGAGAAA CUGAUGA X GAA AGCAUAAA 333 UAUGCUCUU UCUCUUCU AGAAGAGA CUGAUGA X GAA AGAGCAUA 334 AUGCUCUUU CUCUUCUA UAGAAGAG CUGAUGA X GAA AAGAGCAU 335 UGCUCUUUC UCUUCUAC GUAGAAGA CUGAUGA X GAA AAGAGCAU 337 CUCUUUCUC UUCUACAA UUGUAGAA CUGAUGA X GAA AAGAGCAU 339 CUUUCUCUU CUACAAAA UUUUUGAGA CUGAUGA X GAA AAGAAGAG 340 UUUCUCUU UACAAAAA UUUUUGUAC CUGAUGA X GAA AAGAAGAA 342 UCUCUUCUA CAAAAAGC CUUUUGU CUGAUGA X GAA AAGAAGAAA 342 UCUCUUCUA CAAAAGCC GGCUUUUG CUGAUGA X GAA AAGAAGAA 361 UGGAAGAUA AAAUUCGU ACGAAUUU CUGAUGA X GAA AAGAAGAA 366 GAUAAAAUU CGUGAACU AGUUCACG CUGAUGA X GAA AUUUUUAU 367 AUAAAAUU CGUGAACU AGUUCACG CUGAUGA X GAA AUUUUAU 367 AUAAAAUU CGUGAACU GAGUUCAC CUGAUGA X GAA AAUUUUAU 375 CGUGAACUC CGUCCUGA UCAGAACG CUGAUGA X GAA AAUUUUAU 379 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUU 385 GUCCUGAUU GCAUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGAC 386 GAUGACUC UUUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGAC 387 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUU 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUCAGGAC 391 AUUGCAUUU UUUCUGAU AAAAAUGC CUGAUGA X GAA AUCAGGAC 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAUGCAAU 393 UGCAUUUUU UCUGAUAU AUACAGAA CUGAUGA X GAA AAAAAUGCAA 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAAUGCAA 395 CAUUUUUUU CUGAUAUU AAAAAUGC CUGAUGA X GAA AAAAAAUGCAAAAAAAAUGCAAAAAAAAAAAAAAA | | AAAUUUUUU AUGCUCUU | |
| UUUAUGCUC UUUCUCUU AGAAGAA CUGAUGA X GAA AGCAUAAA 333 UAUGCUCUUU UCUCUUCUU AGAAGAGA CUGAUGA X GAA AGAGCAUA 334 AUGCUCUUU CUCUUCUA UAGAAGAG CUGAUGA X GAA AAGAGCAU 335 UGCUCUUUC UCUUCUAC GUAGAAGA CUGAUGA X GAA AAGAGCAU 337 CUCUUUCUC UUCUACAA UUGUAGAA CUGAUGA X GAA AAGAGAGA 339 CUUUCUCUU CUACAAAA UUUUUGUAG CUGAUGA X GAA AGAAAAGAGAAAA 340 UUUCUCUUC UACAAAAA UUUUUGUAG CUGAUGA X GAA AGAAAAAAAAAAAAAAAAAAAAAAAAA | | AAUUUUUUA UGCUCUUU | |
| UAUGCUCUU UCUCUUCU AGAAGAGA CUGAUGA X GAA AGAGCAUA 334 AUGCUCUUU CUCUUCUA UAGAAGAG CUGAUGA X GAA AAGAGCAU 335 UGCUCUUUC UCUUCUAC GUAGAAGA CUGAUGA X GAA AAGAGCAU 337 CUCUUUCUC UUCUACAA UUUUUGAGAA CUGAUGA X GAA AGAAAAGAGA 339 CUUUCUCUU CUACAAAA UUUUUGUAG CUGAUGA X GAA AGAAAAAGAGAAAAAAAAAAAAAAAAA | | | |
| AUGCUCUUU CUCUUCUA UAGAAGAG CUGAUGA X GAA AAGAGCAU 335 UGCUCUUUC UCUUCUAC GUAGAAGA CUGAUGA X GAA AAAGAGCA 337 CUCUUUCUC UUCUACAA UUUUUGAAA CUGAUGA X GAA AGAAAGAG 339 CUUUCUCUU CUACAAAA UUUUUGUAG CUGAUGA X GAA AGAAAAAGAG 340 UUUCUCUUC UACAAAAA UUUUUGUA CUGAUGA X GAA AAGAGAAAA 342 UCUCUUCUA CAAAAGCC GGCUUUUG CUGAUGA X GAA AAAAAAGAGAAA 361 UGGAAGAUA AAAUUCGU ACGAAUUU CUGAUGA X GAA AUCUUCCAA 366 GAUAAAAUU CGUGAACU AGUUCAC CUGAUGA X GAA AUUUUAUCAAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AUUUUAUCAAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AUUUUAUCAAAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AGUUCACCAAAAAAAAAAAAAAAAAAAAAAAAAAA | | UAUGCUCUU UCUCUUCU | |
| UGCUCUUUC UCUUCUAC GUAGAAGA CUGAUGA X GAA AAAGAGCA 337 CUCUUUCUC UUCUACAA UUUUUGAAA CUGAUGA X GAA AGAAAAGAG 339 CUUUCUCUU CUACAAAA UUUUUGUAG CUGAUGA X GAA AGAAAAAGAG 340 UUUCUCUUC UACAAAAG CUUUUUGUA CUGAUGA X GAA AAAAAAAGAGAAAA 342 UCUCUUCUA CAAAAAG CUUUUUGUA CUGAUGA X GAA AAAAAAAGAAAA 361 UGGAAGAUA AAAUUCGU ACGAAUUU CUGAUGA X GAA AUCUUCCA 366 GAUAAAAUU CGUGAACU AGUUCAC CUGAUGA X GAA AUUUUAUCA 367 AUAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AUUUUAUCA 375 CGUGAACUC CGUCCUGA UCAGGACG CUGAUGA X GAA AGUUCACA 379 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUU 385 GUCCUGAUU GCAUUUUU AAAAAUCC CUGAUGA X GAA AUCAGGAC 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAUC 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AUGCAAUC 392 UUGCAUUUU UUCUGAUA UAUCAGAAA CUGAUGA X GAA AAAAAACCAAAAAAAAAAAAAAAAAAAAAAA | | | |
| 337 CUCUUUCUC UUCUACAA 339 CUUUCUCUU CUACAAAA UUUUUGUAG CUGAUGA X GAA AGAGAAAG 340 UUUCUCUUC UACAAAAG CUUUUUGUA CUGAUGA X GAA AAGAGAAA 342 UCUCUUCUA CAAAAGCC GGCUUUUG CUGAUGA X GAA AAGAGAAA 342 UCUCUUCUA CAAAAGCC GGCUUUUG CUGAUGA X GAA AGAGAGAA 361 UGGAAGAUA AAAUUCGU ACGAAUUU CUGAUGA X GAA AUCUUCCA 366 GAUAAAAUU CGUGAACU AGUUCACG CUGAUGA X GAA AUUUUAUC 367 AUAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AAUUUUAUC 375 CGUGAACUC CGUCCUGA UCAGGACG CUGAUGA X GAA AGUUCACC 379 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUU 385 GUCCUGAUU GCAUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGAC 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAUC 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAUC 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAUGCAAUC 393 UGCAUUUUU UCUGAUAU AUAUCAGAA CUGAUGA X GAA AAAAAUGC 394 GCAUUUUUU CUGAUAU AUAUCAGA CUGAUGA X GAA AAAAAUGC 395 CAUUUUUUU UGAUAUG CAUAUCAC CUGAUGA X GAA AAAAAAUGC 396 CAUUUUUUU UCUGAUAUG CAUAUCAC CUGAUGA X GAA AAAAAAUGC 397 CAUUUUUUU CUGAUAUG CAUAUCAC CUGAUGA X GAA AAAAAAUGC 398 CCAUUUUUU CUGAUAUG CAUAUCAC CUGAUGA X GAA AAAAAAUGC 399 CCAUUUUUU CUGAUAUG CAUAUCAC CUGAUGA X GAA AAAAAAUGC 390 CAUUUUUUU CUGAUAUG CAUAUCAC CUGAUGA X GAA AAAAAAUGC 390 CAUUUUUUU CUGAUAUGU ACAUAUCAC CUGAUGA X GAA AAAAAAUGC 390 CAUUUUUUUUUUUU CUGAUAUGU ACAUAUCAC CUGAUGA X GAA AAAAAAUGC 390 CAUUUUUUU CUGAUAUGU ACAUAUCAC CUGAUGA X GAA AAAAAAUGC 390 CAUUUUUUUUUUUUUUUUCUGAU ACAUAUCAC CUGAUGA X GAA AAAAAAUGC 390 CAUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUU | | UGCUCUUUC UCUUCUAC | |
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| 340 UUUCUCUUC UACAAAAG CUUUUGUA CUGAUGA X GAA AAGAGAAA 342 UCUCUUCUA CAAAAGCC GGCUUUUG CUGAUGA X GAA AGAAGAGA 361 UGGAAGAUA AAAUUCGU ACGAAUUU CUGAUGA X GAA AUCUUCCA 366 GAUAAAAUU CGUGAACU AGUUCACG CUGAUGA X GAA AUUUUAU 367 AUAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AAUUUUAU 375 CGUGAACUC CGUCCUGA UCAGGACG CUGAUGA X GAA AGUUCACG 379 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUU 385 GUCCUGAUU GCAUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGAC 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAUC 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAU 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAAGCAA 393 UGCAUUUUU UCUGAUAU AUAUCAGAA CUGAUGA X GAA AAAAAUGC 394 GCAUUUUUU CUGAUAU AUAUCAGA CUGAUGA X GAA AAAAAUGCAA 395 CAUUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAAUGCAA 396 CAUUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAAUGCAAAAAAAUGCAAAAAAAAUGCAAAAAAAAAA | | CUUUCUCUU CUACAAAA | UUUUGUAG CUGAUGA X GAA AGAGAAAG |
| 342 UCUCUUCUA CAAAAGCC GGCUUUUG CUGAUGA X GAA AGAAGAGAAAAAGCG 1961 UGGAAGAUA AAAUUCGU ACGAAUUU CUGAUGA X GAA AUCUUCCAAGA GAAAAAUU CGUGAACU AGUUCACG CUGAUGA X GAA AUUUUAUCGAACU AGUUCACG CUGAUGA X GAA AUUUUAUCGAACUC GAGUUCAC CUGAUGA X GAA AAUUUUAUCGAACUC CGUCCUGA UCAGGACG CUGAUGA X GAA AGUUCACGAACUC CGUCCUGA UCAGGACG CUGAUGA X GAA ACGGAGUUAACGAACUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUUAACAACAACAACAAAAAAAACCAACAACAACAACAACA | | UUUCUCUUC UACAAAAG | CUUUUGUA CUGAUGA X GAA AAGAGAAA |
| 361 UGGAAGAUA AAAUUCGU ACGAAUUU CUGAUGA X GAA AUCUUCCA 366 GAUAAAAUU CGUGAACU AGUUCACG CUGAUGA X GAA AUUUUAUC 367 AUAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AAUUUUAUC 375 CGUGAACUC CGUCCUGA UCAGGACG CUGAUGA X GAA AGUUCACG 379 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUU 385 GUCCUGAUU GCAUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGACG 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAUC 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAUC 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAAUGCAI 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCAI 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUCG 395 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 396 CAUUUUUUU CUGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 397 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 398 CAUUUUUUU CUGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 399 CAUUUUUUU CUGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 390 CAUUUUUUU CUGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 391 CAUUUUUUC CUGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 391 CAUUUUUUU CUGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 391 CAUUUUUUC CUGAUAUGA ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 391 CAUUUUUUCUGAUAUGA ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 391 CAUUUUUCUGAUAUGA ACAUAUCA CUGAUGA X GAA AAAAAAUCGAI 391 CAUUUUUCUGAUAUGA ACAUAUCA CUGAUGA X GAA AAAAAAUCGAIAACGAICGAICGAICGAICGAICGAICGAICGAICG | | UCUCUUCUA CAAAAGCC | GGCUUUUG CUGAUGA X GAA AGAAGAGA |
| 366 GAUAAAAUU CGUGAACU AGUUCACG CUGAUGA X GAA AUUUUAUCGACAC AUAAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AAUUUUAUCGACACACACACACACACACACACACACACAC | | UGGAAGAUA AAAUUCGU | |
| 367 AUAAAAUUC GUGAACUC GAGUUCAC CUGAUGA X GAA AAUUUUAC 375 CGUGAACUC CGUCCUGA UCAGGACG CUGAUGA X GAA AGUUCACG 379 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUU 385 GUCCUGAUU GCAUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGACG 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAUC 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAU 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAAUGCAI 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCAI 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGCAI 395 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUG 396 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUG 397 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 398 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 399 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 390 GCAUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUUC UGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUUC UGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUUC UGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUCUGAUAU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUCUGAUAU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUCUGAUAU AUCAGAA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAUGCAI 391 CAUUUUUCUGAUAU AUCAGAA CUGAUGA X GAA AAAAAAUGCAI 391 CAUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI | | GAUAAAAUU CGUGAACU | AGUUCACG CUGAUGA X GAA AUUUUAUC |
| 375 CGUGAACUC CGUCCUGA UCAGGACG CUGAUGA X GAA AGUUCACG 379 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUU 385 GUCCUGAUU GCAUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGACG 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAUC 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAUC 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAAUGCAA 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCAA 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGCAA 395 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUC 396 CAUUUUUUU UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUC | | AUAAAAUUC GUGAACUC | |
| 379 AACUCCGUC CUGAUUGC GCAAUCAG CUGAUGA X GAA ACGGAGUC 385 GUCCUGAUU GCAUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGAC 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAU 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAU 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAUGCAA 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCA 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGC 395 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAUGCAAUAUCA CUGAUGA X GAA AAAAAAUGCAAUAUCA CUGAUGA X GAA AAAAAAUGCAAUAUCA CUGAUGA X GAA AAAAAAUGCAAUAUCA CUGAUGA X GAA AAAAAAUGCAAUAUCA CUGAUGA X GAA AAAAAAUGCAAUAUCAAUAUCA CUGAUGA X GAA AAAAAAUGCAAUAUCAAUAUCAA CUGAUGA X GAA AAAAAAUGCAAUAUCAAUAUCAA CUGAUGA X GAA AAAAAAAUGCAAUAUCAA CUGAUGA X GAA AAAAAAAUGCAAUAUCAAUAUCAA CUGAUGA X GAA AAAAAAAUGCAAUAUCAAUAUCAA CUGAUGA X GAA AAAAAAAUGCAAUAUCAAUAUCAAUAUCAAUAUCAAUAUCAAUAUCAAAAAA | | CGUGAACUC CGUCCUGA | |
| 385 GUCCUGAUU GCAUUUUU AAAAAUGC CUGAUGA X GAA AUCAGGAC 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAU 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAI 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAUGCAI 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCAI 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGC 395 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUG 396 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 397 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 398 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 399 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 390 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAI 391 AUGCAAUCAGAAA CUGAUGA X GAA AAAAAAUGCAI 392 CAUUUUUUC CUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 393 CAUUUUUUC CUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 394 CAUUUUUUC CUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 395 CAUUUUUUC CUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 395 CAUUUUUUC CUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 396 CAUUUUUUC CUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 397 CAUUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 398 CAUUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 399 CAUUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 390 CAUUUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 390 CAUUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 390 CAUUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 390 CAUUUUUUUCUGAUAUGA CUGAUGA X GAA AAAAAAUGCAI 390 CAUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUUU | | AACUCCGUC CUGAUUGC | GCAAUCAG CUGAUGA X GAA ACGGAGUU |
| 390 GAUUGCAUU UUUUCUGA UCAGAAAA CUGAUGA X GAA AUGCAAUC 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAU 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAUGCAA 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCAA 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGCAA 395 CAUUUUUUC UGAUAUG ACAUAUCA CUGAUGA X GAA AAAAAAUGCAAAAAAAAAAAAAAAAAAAAAAAAAA | | GUCCUGAUU GCAUUUUU | AAAAAUGC CUGAUGA X GAA AUCAGGAC |
| 391 AUUGCAUUU UUUCUGAU AUCAGAAA CUGAUGA X GAA AAUGCAAC 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAUGCAA 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCA 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGCA 395 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUG 396 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAAAAAAAAAAAAAAAAAAAAAAAAAA | | GAUUGCAUU UUUUCUGA | 0 0:10:32.22 |
| 392 UUGCAUUUU UUCUGAUA UAUCAGAA CUGAUGA X GAA AAAUGCAA 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCA 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGCA 395 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCA 396 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCA 397 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCAAAAAAUGCAAAAAAAUGCAAAAAAAAAA | | AUUGCAUUU UUUCUGAU | |
| 393 UGCAUUUUU UCUGAUAU AUAUCAGA CUGAUGA X GAA AAAAUGCA 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGCA 395 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCA 396 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGCA | | UUGCAUUUU UUCUGAUA | UAUCAGAA CUGAUGA X GAA AAAUGCAA |
| 394 GCAUUUUUU CUGAUAUG CAUAUCAG CUGAUGA X GAA AAAAAUGC 395 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGC 396 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUGC | | UGCAUUUUU UCUGAUAU | |
| 395 CAUUUUUUC UGAUAUGU ACAUAUCA CUGAUGA X GAA AAAAAAUC | | GCAUUUUUU CUGAUAUG | |
| CANCIDCA CICALICA Y GAA AUCAGAAA | | CAUUUUUUC UGAUAUGU | ACAUAUCA CUGAUGA X GAA AAAAAAUG |
| 1400 000COGRON COCKECCO COM | 400 | UUUCUGAUA UGUACUUC | GAAGUACA CUGAUGA X GAA AUCAGAAA |

Table III

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| Nt. Position | Substrate | Ribozyme |
| 404 | UGAUAUGUA CUUCCCUU | AAGGGAAG CUGAUGA X GAA ACAUAUCA |
| 404 | UAUGUACUU CCCUUGGA | UCCAAGGG CUGAUGA X GAA AGUACAUA |
| | AUGUACUUC CCUUGGAC | GUCCAAGG CUGAUGA X GAA AAGUACAU |
| 408 | ACUUCCCUU GGACAGUA | UACUGUCC CUGAUGA X GAA AGGGAAGU |
| 412 | UGGACAGUA GAUAUUGC | GCAAUAUC CUGAUGA X GAA ACUGUCCA |
| 420 | CAGUAGAUA UUGCUGAU | AUCAGCAA CUGAUGA X GAA AUCUACUG |
| 424 | | UCAUCAGC CUGAUGA X GAA AUAUCUAC |
| 426 | COLLECTION | GGGAUGUG CUGAUGA X GAA AGCUCAUC |
| 438 | | AGGGAUGU CUGAUGA X GAA AAGCUCAU |
| 439 | AUGAGCUUC ACAUCCCU | AUACGAGG CUGAUGA X GAA AUGUGAAG |
| 444 | CUUCACAUC CCUCGUAU | CAAAAUAC CUGAUGA X GAA AGGGAUGU |
| 448 | ACAUCCCUC GUAUUUUG | GUACAAAA CUGAUGA X GAA ACGAGGGA |
| 451 | UCCCUCGUA UUUUGUAC | 001101222 |
| 453 | CCUCGUAUU UUGUACAA | 0000110121 000110111 |
| 454 | CUCGUAUUU UGUACAAU | |
| 455 | UCGUAUUUU GUACAAUU | 7210000110 00011011 |
| 458 | UAUUUUGUA CAAUUUGU | 110/HH1000 OUG110011 |
| 463 | UGUACAAUU UGUCUGCU | 7.007.01.01 |
| 464 | GUACAAUUU GUCUGCUU | |
| 467 | CAAUUUGUC UGCUUACA | UGUAAGCA CUGAUGA X GAA ACAAAUUG |
| 472 | UGUCUGCUU ACAUGUGC | GCACAUGU CUGAUGA X GAA AGCAGACA |
| 473 | GUCUGCUUA CAUGUGCU | AGCACAUG CUGAUGA X GAA AAGCAGAC |
| 482 | CAUGUGCUA CAGCAUUA | UAAUGCUG CUGAUGA X GAA AGCACAUG |
| 489 | UACAGCAUU AUGCACAA | |
| 490 | ACAGCAUUA UGCACAAC | GUUGUGCA CUGAUGA X GAA AAUGCUGU |
| 501 | CACAACCUU AAGGUUUA | UAAACCUU CUGAUGA X GAA AGGUUGUG |
| 502 | ACAACCUUA AGGUUUAC | GUAAACCU CUGAUGA X GAA AAGGUUGU |
| 507 | CUUAAGGUU UACAGACC | GGUCUGUA CUGAUGA X GAA ACCUUAAG |
| 508 | UUAAGGUUU ACAGACCU | AGGUCUGU CUGAUGA X GAA AACCUUAA |
| 509 | UAAGGUUUA CAGACCUC | GAGGUCUG CUGAUGA X GAA AAACCUUA |
| 517 | ACAGACCUC ACAAGCAG | CUGCUUGU CUGAUGA X GAA AGGUCUGU |
| 529 | AGCAGCCUA AUCUAGAC | GUCUAGAU CUGAUGA X GAA AGGCUGCU |
| 532 | AGCCUAAUC UAGACGAA | UUCGUCUA CUGAUGA X GAA AUUAGGCU |
| 534 | CCUAAUCUA GACGAAUC | GAUUCGUC CUGAUGA X GAA AGAUUAGG |
| 542 | AGACGAAUC UCAAAGUU | AACUUUGA CUGAUGA X GAA AUUCGUCU |
| 544 | ACGAAUCUC AAAGUUUC | GAAACUUU CUGAUGA X GAA AGAUUCGU |
| 550 | CUCAAAGUU UCGUGGUU | AACCACGA CUGAUGA X GAA ACUUUGAG |
| 551 | UCAAAGUUU CGUGGUUC | GAACCACG CUGAUGA X GAA AACUUUGA |
| 552 | CAAAGUUUC GUGGUUCC | GGAACCAC CUGAUGA X GAA AAACUUUG |
| 558 | UUCGUGGUU CCUGGUUU | AAACCAGG CUGAUGA X GAA ACCACGAA |
| 559 | UCGUGGUUC CUGGUUUA | UAAACCAG CUGAUGA X GAA AACCACGA |
| 565 | UUCCUGGUU UACCUGAU | AUCAGGUA CUGAUGA X GAA ACCAGGAA |
| 566 | UCCUGGUUU ACCUGAUG | CAUCAGGU CUGAUGA X GAA AACCAGGA |
| 567 | CCUGGUUUA CCUGAUGA | UCAUCAGG CUGAUGA X GAA AAACCAGG |
| 579 | GAUGAGAUA AAGUUCAA | UUGAACUU CUGAUGA X GAA AUCUCAUC |
| 584 | GAUAAAGUU CAAGUUAU | AUAACUUG CUGAUGA X GAA ACUUUAUC |
| 585 | AUAAAGUUC AAGUUAUC | GAUAACUU CUGAUGA X GAA AACUUUAU |
| 590 | GUUCAAGUU AUCCCAAC | GUUGGGAU CUGAUGA X GAA ACUUGAAC |
| 591 | UUCAAGUUA UCCCAACU | AGUUGGGA CUGAUGA X GAA AACUUGAA |
| 593 | CAAGUUAUC CCAACUGA | UCAGUUGG CUGAUGA X GAA AUAACUUG |
| 610 | CAGAUGAUC UGAGAAAG | CUUUCUCA CUGAUGA X GAA AUCAUCUG |
| 620 | GAGAAAGUC GGAUGACC | GGUCAUCC CUGAUGA X GAA ACUUUCUC |
| 639 | AAGACUGUU UUUGACGA | UCGUCAAA CUGAUGA X GAA ACAGUCUU |
| .640 | AGACUGUUU UUGACGAA | UUCGUCAA CUGAUGA X GAA AACAGUCU |
| 641 | GACUGUUUU UGACGAAU | AUUCGUCA CUGAUGA X GAA AAACAGUC |
| 642 | ACUGUUUUU GACGAAUU | AAUUCGUC CUGAUGA X GAA AAAACAGU |
| 650 | UGACGAAUU GCUCGAAC | GUUCGAGC CUGAUGA X GAA AUUCGUCA |
| 654 | GAAUUGCUC GAACAAGU | ACUUGUUC CUGAUGA X GAA AGCAAUUC |
| . 003 | | |

Table III

| Nt. Position | Substrate | Ribozyme |
|-----------------|------------------------------|------------------------------------|
| 663 | GAACAAGUU GAAGAUUC | GAAUCUUC CUGAUGA X GAA ACUUGUUC |
| 670 | UUGAAGAUU CGGAGGAA | UUCCUCCG CUGAUGA X GAA AUCUUCAA |
| 671 | UGAAGAUUC GGAGGAAC | GUUCCUCC CUGAUGA X GAA AAUCUUCA |
| 686 | ACGAAGCUA UGGCAUUG | CAAUGCCA CUGAUGA X GAA AGCUUCGU |
| 693 | UAUGGCAUU GUUCAUGA | UCAUGAAC CUGAUGA X GAA AUGCCAUA |
| | GGCAUUGUU CAUGAUAC | GUAUCAUG CUGAUGA X GAA ACAAUGCC |
| 696 697 | GCAUUGUUC AUGAUACA | UGUAUCAU CUGAUGA X GAA AACAAUGC |
| 703 | UUCAUGAUA CAUUUUAU | AUAAAAUG CUGAUGA X GAA AUCAUGAA |
| 707 | UGAUACAUU UUAUGAGC | GCUCAUAA CUGAUGA X GAA AUGUAUCA |
| 708 | GAUACAUUU UAUGAGCU | AGCUCAUA CUGAUGA X GAA AAUGUAUC |
| 709 | AUACAUUUU AUGAGCUA | UAGCUCAU CUGAUGA X GAA AAAUGUAU |
| 710 | UACAUUUUA UGAGCUAG | CUAGCUCA CUGAUGA X GAA AAAAUGUA |
| 717 | UAUGAGCUA GAACCUGC | GCAGGUUC CUGAUGA X GAA AGCUCAUA |
| 728 | ACCUGCAUA UGUUGACU | AGUCAACA CUGAUGA X GAA AUGCAGGU |
| 732 | GCAUAUGUU GACUACUA | UAGUAGUC CUGAUGA X GAA ACAUAUGC |
| 737 | UGUUGACUA CUACCAGA | UCUGGUAG CUGAUGA X GAA AGUCAACA |
| 740 | UGACUACUA CCAGAAAU | AUUUCUGG CUGAUGA X GAA AGUAGUCA |
| 749 | CCAGAAAUU AAAGAAAC | GUUUCUUU CUGAUGA X GAA AUUUCUGG |
| 750 | CAGAAAUUA AAGAAACC | GGUUUCUU CUGAUGA X GAA AAUUUCUG |
| 766 | CAAAAUGUU GGCAUUUU | AAAAUGCC CUGAUGA X GAA ACAUUUUG |
| 772 | GUUGGCAUU UUGGUCCG | CGGAO~AA CUGAUGA X GAA AUGCCAAC |
| 773 | UUGGCAUUU UGGUCCGC | GCGGACCA CUGAUGA X GAA AAUGCCAA |
| 774 | UGGCAUUUU GGUCCGCU | AGCGGACC CUGAUGA X GAA AAAUGCCA |
| 778 | AUUUUGGUC CGCUCUCU | AGAGAGCG CUGAUGA A GAA ACCAAAAU |
| 783 | GGUCCGCUC UCUCAUUU | AAAUGAGA CUGAUGA X GAA AGCGGACC |
| 785 | UCCGCUCUC UCAUUUUG | CAAAAUGA CUGAUGA X GAA AGAGCGGA |
| 787 | CGCUCUCUC AUUUUGCA | UGCAAAAU CUGAUGA X GAA AGAGAGCG |
| 790 | UCUCUCAUU UUGCAUCC | GGAUGCAA CUGAUGA X GAA AUGAGAGA |
| 791 | CUCUCAUUU UGCAUCCA | UGGAUGCA CUGAUGA X GAA AAUGAGAG |
| 792 | UCUCAUUUU GCAUCCAA | UUGGAUGC CUGAUGA X GAA AAAUGAGA |
| 797 | UUUUGCAUC CAAAUCCG | CGGAUUUG CUGAUGA X GAA AUGCAAAA |
| 803 | AUCCAAAUC CGUAGUAA | UUACUACG CUGAUGA X GAA AUUUGGAU |
| 807 | AAAUCCGUA GUAAGGAA | UUCCUUAC CUGAUGA X GAA ACGGAUUU |
| 810 | UCCGUAGUA AGGAACUA | UAGUUCCU CUGAUGA X GAA ACUACGGA |
| 818 | AAGGAACUA AUUUCUGA | UCAGAAAU CUGAUGA X GAA AGUUCCUU |
| 821 | GAACUAAUU UCUGAGCA | UGCUCAGA CUGAUGA X GAA AUUAGUUC |
| 822 | AACUAAUUU CUGAGCAU | AUGCUCAG CUGAUGA X GAA AAUUAGUU |
| 823 | ACUAAUUUC UGAGCAUA | UAUGCUCA CUGAUGA X GAA AAAUUAGU |
| 831 | CUGAGCAUA ACAACAAU | AUUGUUGU CUGAUGA X GAA AUGCUCAG |
| 845 | AAUGAGAUU GUUAUAGA | UCUAUAAC CUGAUGA X GAA AUCUCAUU |
| 848 | GAGAUUGUU AUAGAUUG | CAAUCUAU CUGAUGA X GAA ACAAUCUC |
| 849 | AGAUUGUUA UAGAUUGG | CCAAUCUA CUGAUGA X GAA AACAAUCU |
| 851 | AUUGUUAUA GAUUGGUU | AACCAAUC CUGAUGA X GAA AUAACAAU |
| 855 | UUAUAGAUU GGUUGAAU | AUUCAACC CUGAUGA X GAA AUCUAUAA |
| 859 | AGAUUGGUU GAAUGCAC | GUGCAUUC CUGAUGA X GAA ACCAAUCU |
| 876 | AGAAACCUA AAUCGGUU | AACCGAUU CUGAUGA X GAA AGGUUUCU |
| 880 | ACCUAAAUC GGUUCUCU | AGAGAACC CUGAUGA X GAA AUUUAGGU |
| 884 | AAAUCGGUU CUCUAUGU | ACAUAGAG CUGAUGA X GAA ACCGAUUU |
| 885 | AAUCGGUUC UCUAUGUA | UACAUAGA CUGAUGA X GAA AACCGAUU |
| 887 | UCGGUUCUC UAUGUAUC | GAUACAUA CUGAUGA X GAA AGAACCGA |
| 889 | GGUUCUCUA UGUAUCUU | AAGAUACA CUGAUGA X GAA AGAGAACC |
| 893 | CUCUAUGUA UCUUUCGG | CCGAAAGA CUGAUGA X GAA ACAUAGAG |
| 895 | CUAUGUAUC UUUCGGAA | UUCCGAAA CUGAUGA X GAA AUACAUAG |
| 897 | AUGUAUCUU UCGGAAGC | GCUUCCGA CUGAUGA X GAA AGAUACAU |
| | UGUAUCUUU CGGAAGCA | UGCUUCCG CUGAUGA X GAA AAGAUACA |
| 898 | I I I (SI) A U CUUU CUMAAAAA | UGCUUCCG CUGAUGI II GIZI IZIGIIGII |

Table III

| | | | Lavin |
|--|----------|--|---------------------------------|
| | Nt. | Substrate | Ribozyme |
| | Position | | AGGAAAUC CUGAUGA X GAA AGCCAUGC |
| • | 912 | GCAUGGCUA GAUUUCCU | UCUCAGGA CUGAUGA X GAA AUCUAGCC |
| | 916 | GGCUAGAUU UCCUGAGA | CUCUCAGG CUGAUGA X GAA AAUCUAGC |
| | 917 | GCUAGAUUU CCUGAGAG | GCUCUCAG CUGAUGA X GAA AAAUCUAG |
| • | 918 | CUAGAUUUC CUGAGAGC | GCUUGGGC CUGAUGA X GAA AUUUCAUU |
| | 941 | AAUGAAAUA GCCCAAGC | AGCAUCCA CUGAUGA X GAA AGCUUGGG |
| | 951 | CCCAAGCUC UGGAUGCU | AACAUUUG CUGAUGA X GAA AGCAUCCA |
| • | 960 | UGGAUGCUU CAAAUGUU | GAACAUUU CUGAUGA X GAA AAGCAUCC |
| | 961 | GGAUGCUUC AAAUGUUC | AUGAAAGG CUGAUGA X GAA ACAUUUGA |
| | 968 | UCAAAUGUU CCUUUCAU | AAUGAAAG CUGAUGA X GAA AACAUUUG |
| | 969 | CAAAUGUUC CUUUCAUU | AAAAAUGA CUGAUGA X GAA AGGAACAU |
| | 972 | AUGUUCCUU UCAUUUUU | CAAAAAUG CUGAUGA X GAA AAGGAACA |
| | 973 | UGUUCCUUU CAUUUUUG | ACAAAAAU CUGAUGA X GAA AAAGGAAC |
| | 974 | GUUCCUUUC AUUUUUGU | AAUACAAA CUGAUGA X GAA AUGAAAGG |
| | 977 | CCUUUCAUU UUUGUAUU | CAAUACAA CUGAUGA X GAA AAUGAAAG |
| 1 (1) | 978 | CUUUCAUUU UUGUAUUG | UCAAUACA CUGAUGA X GAA AAAUGAAA |
| | 979 | UUUCAUUUU UGUAUUGA | CUCAAUAC CUGAUGA X GAA AAAAUGAA |
| | 980 | UUCAUUUUU GUAUUGAG | GGCCUCAA CUGAUGA X GAA ACAAAAAU |
| | 983 | AUULJUUGUA UUGAGGCC | UAGGCCUC CUGAUGA X GAA AUACAAAA |
| | 985 | UUUUGUAUU GAGGCCUA | UUCUUCAU CUGAUGA X GAA AGGCCUCA |
| | 993 | UGAGGCCUA AUGAAGAA | GCAACCAC CUGAUGA X GAA ACGCCGUU |
| and the september of the second | 1009 | AACGGCGUC GUGGUUGC | CAACUGGC CUGAUGA X GAA ACCACGAC |
| | 1015 | GUCGUGGUU GCCAGUUG | AAAUUACC CUGAUGA X GAA ACUGGCAA |
| المراجع المتعالم المت | 1022 | UUGCCAGUU GGUAAUUU | CUCUAAAU CUGAUGA X GAA ACCAACUG |
| | 1026 | CAGUUGGUA AUUUAGAG | GUCCUCUA CUGAUGA X GAA AUUACCAA |
| | 1029 | UUGGUAAUU UAGAGGAC UGGUAAUUU AGAGGACA | UGUCCUCU CUGAUGA X GAA AAUUACCA |
| | 1030 | GGUAAUUUA GAGGACAA | UUGUCCUC CUGAUGA X GAA AAAUUACC |
| | 1031 | ACAAGACUA AAAAGGGU | ACCCUUUU CUGAUGA X GAA AGUCUUGU |
| 4 - 4 - 4 | 1044 | AAAAGGGUU UGUACAUC | GAUGUACA CUGAUGA X GAA ACCCUUUU |
| | 1053 | AAAGGGUUU GUACAUCA | UGAUGUAC CUGAUGA X GAA AACCCUUU |
| E trata | 1054 | GGGUUUGUA CAUCAAAG | CUUUGAUG CUGAUGA X GAA ACAAACCC |
| | 1057 | UUGUACAUC AAAGGGUG | CACCCUUU CUGAUGA X GAA AUGUACAA |
| المرائح يجازو مبلغ بسجانها | 1061 | GGGUGGGUC CCACAGCU | AGCUGUGG CUGAUGA X GAA ACCCACCC |
| * | 1073 | CCACAGCUU ACGAUCAU | AUGAUCGU CUGAUGA X GAA AGCUGUGG |
| • | 1082 | CACAGCUUA CGAUCAUG | CAUGAUCG CUGAUGA X GAA AAGCUGUG |
| | 1088 | CUUACGAUC AUGGAACA | UGUUCCAU CUGAUGA X GAA AUCGUAAG |
| | 1098 | UGGAACAUU CAGCAACA | UGUUGCUG CUGAUGA X GAA AUGUUCCA |
| • | 1099 | GGAACAUUC AGCAACAG | CUGUUGCU CUGAUGA X GAA AAUGUUCC |
| | 1114 | AGGCGGGUU CAUGACUC | GAGUCAUG CUGAUGA X GAA ACCCGCCU |
| | 1115 | GGCGGGUUC AUGACUCA | UGAGUCAU CUGAUGA X GAA AACCCGCC |
| | 1122 | UCAUGACUC AUUGUGGU | ACCACAAU CUGAUGA X GAA AGUCAUGA |
| | 1125 | UGACUCAUU GUGGUACU | AGUACCAC CUGAUGA X GAA AUGAGUCA |
| | 1131 | AUUGUGGUA CUAAUUCG | CGAAUUAG CUGAUGA X GAA ACCACAAU |
| • | 1134 | GUGGUACUA AUUCGGUU | AACCGAAU CUGAUGA X GAA AGUACCAC |
| • | 1137 | GUACUAAUU CGGTJUCUG | CAGAACCG CUGAUGA X GAA AUUAGUAC |
| | 1138 | UACUAAUUC GGUUCUGG | CCAGAACC CUGAUGA X GAA AAUUAGUA |
| | 1142 | AAUUCGGUU CUGGAAGC | GCUUCCAG CUGAUGA X GAA ACCGAAUU |
| | 1143 | AUUCGGUUC UGGAAGCC | GGCUUCCA CUGAUGA X GAA AACCGAAU |
| | 1154 | GAAGCCAUC ACUUUUGG | CCAAAAGU CUGAUGA X GAA AUGGCUUC |
| | 1158 | CCAUCACUU UUGGCGUG | CACGCCAA CUGAUGA X GAA AGUGAUGG |
| - | 1159 | CAUCACUUU UGGCGUGC | GCACGCCA CUGAUGA X GAA AAGUGAUO |
| | 1160 | AUCACUUUU GGCGUGCC | GGCACGCC CUGAUGA X GAA AAAGUGAU |
| • | 1175 | CCAAUGAUA ACAUGGCC | GGCCAUGU CUGAUGA X GAA AUCAUUGG |
| | 1187 | UGGCCACUU UAUGCUGA | UCAGCAUA CUGAUGA X GAA AGUGGCCA |
| | 1188 | GGCCACUUU AUGCUGAU | AUCAGCAU CUGAUGA X GAA AAGUGGCC |
| | 1189 | GCCACUUUA UGCUGAUC | GAUCAGCA CUGAUGA X GAA AAAGUGGC |
| • | 1197 | AUGCUGAUC AAUUCUAC | GUAGAAUU CUGAUGA X GAA AUCAGCAU |
| | , | | |

Table III

| | | I Débassion |
|----------|---------------------|--|
| Nt. | Substrate | Ribozyme |
| Position | TICTUCA ATTUCATOR | CGUUGUAG CUGAUGA X GAA AUUGAUCA |
| 1201 | UGAUCAAUU CUACAACG | UCGUUGUA CUGAUGA X GAA AAUUGAUC |
| 1202 | GAUCAAUUC UACAACGA | UCUCGUUG CUGAUGA X GAA AGAAUUGA |
| 1204 | UCAAUUCUA CAACGAGA | ACCUCGAC CUGAUGA X GAA ACCUUCUC |
| 1217 | GAGAAGGUA GUCGAGGU | CUAACCUC CUGAUGA X GAA ACCOCCU |
| 1220 | AAGGUAGUC GAGGUUAG | |
| 1226 | GUCGAGGUU AGGGGAUU | |
| 1227 | UCGAGGUUA GGGGAUUG | CAAUCCCC CUGAUGA X GAA AACCUCGA |
| 1234 | UAGGGGAUU GGGAAUCA | UGAUUCCC CUGAUGA X GAA AUCCCCUA |
| 1241 | UUGGGAAUC AAAAUCGG | CCGAUUUU CUGAUGA X GAA AUUCCCAA |
| 1247 | AUCAAAAUC GGGAUAGA | UCUAUCCC CUGAUGA X GAA AUUUUGAU |
| 1253 | AUCGGGAUA GAUGUAUG | CAUACAUC CUGAUGA X GAA AUCCCGAU |
| 1259 | AUAGAUGUA UGGAAUGA | UCAUUCCA CUGAUGA X GAA ACAUCUAU |
| 1274 | GAAGGGAUU GAGAUCAC | GUGAUCUC CUGAUGA X GAA AUCCCUUC |
| 1280 | AUUGAGAUC ACGGGCCC | GGGCCCGU CUGAUGA X GAA AUCUCAAU |
| 1292 | GGCCCUGUA AUAGAAAG | CUUUCUAU CUGAUGA X GAA ACAGGGCC |
| 1295 | CCUGUAAUA GAAAGCGC | GCGCUUUC CUGAUGA X GAA AUUACAGG |
| 1310 | GCCAAGAUU AGAGAAGC | GCUUCUCU CUGAUGA X GAA AUCUUGGC |
| 1311 | CCAAGAUUA GAGAAGCA | UGCUUCUC CUGAUGA X GAA AAUCUUGG |
| 1322 | GAAGCAAUU GAGAGACU | AGUCUCUC CUGAUGA X GAA AUUGCUUC |
| 1331 | GAGAGACUA AUGAUCAG | CUGAUCAU CUGAUGA X GAA AGUCUCUC |
| 1337 | CUAAUGAUC AGUAAUGG | CCAUUACU CUGAUGA X GAA AUCAUUAG |
| 1341 | UGAUCAGUA AUGGUUCU | AGAACCAU CUGAUGA X GAA ACUGAUCA |
| 1347 | GUAAUGGUU CUGAGGAA | UUCCUCAG CUGAUGA X GAA ACCAUUAC |
| 1348 | UAAUGGUUC UGAGGAAA | UUUCCUCA CUGAUGA X GAA AACCAUUA |
| | GAGGAAAUU AUAAAUAU | AUAUUUAU CUGAUGA X GAA AUUUCCUC |
| 1358 | AGGAAAUUA UAAAUAUU | AAUAUUUA CUGAUGA X GAA AAUUUCCU |
| 1359 | GAAAUUAUA AAUAUUAG | CUAAUAUU CUGAUGA X GAA AUAAUUUC |
| 1361 | UUAUAAAUA UUAGGGAU. | AUCCCUAA CUGAUGA X GAA AUUUAUAA |
| 1365 | AUAAAUAUU AGGGAUAG | CUAUCCCU CUGAUGA X GAA AUAUUUAU |
| 1367 | UAAAUAUUA GGGAUAGA | UCUAUCCC CUGAUGA X GAA AAUAUUUA |
| 1368 | UAAAUAUUA GOGAUAGA | CAUUACUC CUGAUGA X GAA AUCCCUAA |
| 1374 | UUAGGGAUA GAGUAAUG | AUAGCCAU CUGAUGA X GAA ACUCUAUC |
| 1379 | GAUAGAGUA AUGGCUAU | UUUGCUCA CUGAUGA X GAA AGCCAUUA |
| 1386 | UAAUGGCUA UGAGCAAA | UGCAUUCU CUGAUGA X GAA AGCCAUUU |
| 1401 | AAAUGGCUC AGAAUGCA | UCCACGAA CUGAUGA X GAA AUCCACCU |
| 1426 | AGGUGGAUC UUCGUGGA | GUUCCACG CUGAUGA X GAA AGAUCCAC |
| 1428 | GUGGAUCUU CGUGGAAC | UGUUCCAC CUGAUGA X GAA AAGAUCCA |
| 1429 | UGGAUCUUC GUGGAACA | AGCAGUGA CUGAUGA X GAA AUUGUUCC |
| 1440 | GGAACAAUC UCACUGCU | AGAGCAGU CUGAUGA X GAA AGAUUGUU |
| 1442 | AACAAUCUC ACUGCUCU | UUGAAUGA CUGAUGA X GAA AGCAGUGA |
| 1449 | UCACUGCUC UCAUUCAA | UGUUGAAU CUGAUGA X GAA AGCAGUGA UGUUGAAU CUGAUGA X GAA AGAGCAGU |
| 1451 | ACUGCUCUC AUUCAACA | AUAUGUUG CUGAUGA X GAA AUGAGAGC |
| 1454 | GCUCUCAUU CAACAUAU | GAUAUGUU CUGAUGA X GAA AAUGAGAG |
| 1455 | CUCUCAUUC AACAUAUC | AUUCUUGA CUGAUGA X GAA AUGUUGAA |
| 1461 | UUCAACAUA UCAAGAAU | HADDUCTURA CUCAUCA V CAR AUGUOGAA |
| 1463 | CAACAUAUC AAGAAUUA | UAAUUCUU CUGAUGA X GAA AUAUGUUG |
| 1470 | UCAAGAAUU AUAAUCUU | AAGAUUAU CUGAUGA X GAA AUUCUUGA UAAGAUUA CUGAUGA X GAA AAUUCUUG |
| 1471 | CAAGAAUUA UAAUCUUA | |
| 1473 | AGAAUUAUA AUCUUAAU | AUUAAGAU CUGAUGA X GAA AUAAUUCU |
| 1476 | AUUAUAAUC UUAAUUAG | CUAAUUAA CUGAUGA X GAA AUUAUAAU |
| 1478 | UAUAAUCUU AAUUAGUU | AACUAAUU CUGAUGA X GAA AGAUUAUA |
| 1479 | AUAAUCUUA AUUAGUUG | CAACUAAU CUGAUGA X GAA AAGAUUAU |
| 1482 | AUCUUAAUU AGUUGAAG | CUUCAACU CUGAUGA X GAA AUUAAGAU |
| 1483 | UCUUAAUUA GUUGAAGA | UCUUCAAC CUGAUGA X GAA AAUUAAGA |
| 1486 | UAAUUAGUU GAAGACAG | CUGUCUUC CUGAUGA X GAA ACUAAUUA |
| 1499 | ACAGAAAUA AGUCCUUG | CAAGGACU CUGAUGA X GAA AUUUCUGU |
| 1503 | AAAUAAGUC CUUGCAUU | AAUGCAAG CUGAUGA X GAA ACUUAUUU |
| 1506 | UAAGUCCUU GCAUUGUA | UACAAUGC CUGAUGA X GAA AGGACUUA |
| | 1 | |

Table III

| Nt. | Substrate | Ribozyme |
|----------|--------------------|---------------------------------|
| Position | · | W GDD DUGGDDGG |
| 1511 | CCUUGCAUU GUAACAUG | CAUGUUAC CUGAUGA X GAA AUGCAAGG |
| 1514 | UGCAUUGUA ACAUGGUG | CACCAUGU CUGAUGA X GAA ACAAUGCA |
| 1534 | GUGUGUGUU UUUUUUCC | GGAAAAA CUGAUGA X GAA ACACACAC |
| 1535 | UGUGUGUUU UUUUUCCA | UGGAAAAA CUGAUGA X GAA AACACACA |
| 1536 | GUGUGUUUU UUUUCCAC | GUGGAAAA CUGAUGA X GAA AAACACAC |
| 1537 | UGUGUUUUU UUUCCACU | AGUGGAAA CUGAUGA X GAA AAAACACA |
| 1538 | GUGUUUUUU UUCCACUU | AAGUGGAA CUGAUGA X GAA AAAAACAC |
| 1539 | UGUUUUUUU UCCACUUA | UAAGUGGA CUGAUGA X GAA AAAAAACA |
| 1540 | GUUUUUUU CCACUUAA | UUAAGUGG CUGAUGA X GAA AAAAAAAC |
| 1541 | UUUUUUUUC CACUUAAU | AUUAAGUG CUGAUGA X GAA AAAAAAA |
| 1546 | UUUCCACUU AAUAAAAU | AUUUUAUU CUGAUGA X GAA AGUGGAAA |
| 1547 | UUCCACUUA AUAAAAUG | CAUUUUAU CUGAUGA X GAA AAGUGGAA |
| 1550 | CACUUAAUA AAAUGAAG | CUUCAUUU CUGAUGA X GAA AUUAAGUG |
| 1579 | GGAUGGAUC UUAACUUU | AAAGUUAA CUGAUGA X GAA AUCCAUCC |
| 1581 | AUGGAUCUU AACUUUAA | UUAAAGUU CUGAUGA X GAA AGAUCCAU |
| 1582 | UGGAUCUUA ACUUUAAA | UUUAAAGU CUGAUGA X GAA AAGAUCCA |
| 1586 | UCUUAACUU UAAAAAA | UUUUUUUA CUGAUGA X GAA AGUUAAGA |
| 1587 | CUUAACUUU AAAAAAAA | UUUUUUUU CUGAUGA X GAA AAGUUAAG |
| 1588 | UUAACUUUA AAAAAAAA | UUUUUUUU CUGAUGA X GAA AAAGUUAA |

Where "X" represents stem II region of a HH ribozyme (Hertel et al., 1992 <u>Nucleic Acids Res.</u> 20 3252). The length of stem II may be \geq 2 base-pairs.

Table IV: Solanidine glucosyltransferase Hairpin
Ribozyme and Target Sequences

| 79 | AUGACC AGAA GAUA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UAUCC GCU GGUCAU |
|------|---|------------------|
| 211 | UGGGAA AGAA GGAA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UUCCG GAU UUCCCA |
| 249 | AACUUC AGAA GAGG ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | CCUCU GCU GAAGUU |
| 376 | AAUCAG AGAA GAGU ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | ACUCC GUC CUGAUU |
| 381 | AAUGCA AGAA GGAC ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | GUCCU GAU UGCAUU |
| 429 | AAGCUC AGAA GCAA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UUGCU GAU GAGCUU |
| 468 | CAUGUA AGAA GACA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UGUCU GCU UACAUG |
| 511 | UGUGAG AGAA GUAA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UUACA GAC CUCACA |
| 524 | AGAUUA AGAA GCUU ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | AAGCA GCC UAAUCU |
| 570 | HAUCUC AGAA GGUA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UACCU GAU GAGAUA |
| 603 | CAGAUC AGAA GUCA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UGACA GAU GAUCUG |
| 621 | UUGGUC AGAA GACU ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | AGUCG GAU GACCAA |
| 636 | GUCAAA AGAA GUCU ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | AGACU GUU UUUGAC |
| 779 | UGAGAG AGAA GACC ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | GGUCC GCU CUCUCA |
| 881 | AUAGAG AGAA GAUU ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | AAUCG GUU CUCUAU |
| 1019 | AUUACC AGAA GGCA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UGCCA GUU GGUAAU |
| 1078 | AUCGUA AGAA GUGG ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | CCACA GCU UACGAU |
| 1139 | UUCCAG AGAA GAAU ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | AUUCG GUU CUGGAA |
| 1193 | GAAUUG AGAA GCAU ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | AUGCU GAU CAAUUC |
| 1445 | AAUGAG AGAA GUGA ACCAGAGAAACACACGUUGUGGUACAUUACCUGGUA | UCACU GCU CUCAUU |
| 1933 | 1210010 11011 01011 | |

Table V

Table V: Potato Citrate Synthase Hammerhead Ribozyme and Target Sequences

| | | Ribozyme |
|----------|--------------------|---------------------------------|
| Nt. | Substrate | RIDOZYMe |
| Position | 20200 | GCUGAUGG CUGAUGA X GAA ACGAAAAA |
| 9 | UUUUUCGUU CCAUCAGC | GCUGAUGC CUGAUGA X GAA ACGAAAAA |
| 10 | UUUUCGUUC CAUCAGCC | 00000110 |
| 14 | CGUUCCAUC AGCCUACU | 110011000 |
| 20 | AUCAGCCUA CUUGAGAU | |
| 23 | AGCCUACUU GAGAUGUA | CACHECOC COLLEGE |
| 31 | UGAGAUGUA UUCCCACU | AGUGGGAA CUGAUGA X GAA ACAUCUCA |
| 33 | AGAUGUAUU CCCACUGG | CCAGUGGG CUGAUGA X GAA AUACAUCU |
| 34 | GAUGUAUUC CCACUGGU | ACCAGUGG CUGAUGA X GAA AAUACAUC |
| 43 | CCACUGGUA AAAGUUAA | UUAACUUU CUGAUGA X GAA ACCAGUGG |
| 49 | GUAAAAGUU AAUUUUUU | AAAAAAUU CUGAUGA X GAA ACUUUUAC |
| 50 | UAAAAGUUA AUUUUUUU | AAAAAAU CUGAUGA X GAA AACUUUUA |
| 53 | AAGUUAAUU UUUUUGAU | AUCAAAAA CUGAUGA X GAA AUUAACUU |
| 54 | AGUUAAUUU UUUUGAUU | AAUCAAAA CUGAUGA X GAA AAUUAACU |
| 55 | GUUAAUUUU UUUGAUUU | AAAUCAAA CUGAUGA X GAA AAAUUAAC |
| .56 | UUAAUUUUU UUGAUUUU | AAAAUCAA CUGAUGA X GAA AAAAUUAA |
| 5.7 | UAAUUUUU UGAUUUUC | GAAAAUCA CUGAUGA X GAA AAAAAUUA |
| 58 | AAUUUUUUU GAUUUUCG | CGAAAAUC CUGAUGA X GAA AAAAAAUU |
| 62 | UUUUUGAUU UUCGCGAG | CUCGCGAA CUGAUGA X GAA AUCAAAAA |
| 63 | UUUUGAUUU UCGCGAGC | GCUCGCGA CUGAUGA X GAA AAUCAAAA |
| 64 | UUUGAUUUU CGCGAGCA | UGCUCGCG CUGAUGA X GAA AAAUCAAA |
| 65 | UUGAUUUUC GCGAGCAA | UUGCUCGC CUGAUGA X GAA AAAAUCAA |
| 80 | AAUGGUGUU CUACCGUA | UACGGUAG CUGAUGA X GAA ACACCAUU |
| 81 | AUGGUGUUC UACCGUAG | CUACGGUA CUGAUGA X GAA AACACCAU |
| 83 | GGUGUUCUA CCGUAGCG | CGCUACGG CUGAUGA X GAA AGAACACC |
| 88 | UCUACCGUA GCGUUUCG | CGAAACGC CUGAUGA X GAA ACGGUAGA |
| 9.3 | CGUAGCGUU UCGUUGCU | AGCAACGA CUGAUGA X GAA ACGCUACG |
| 94 | GUAGCGUUU CGUUGCUG | CAGCAACG CUGAUGA X GAA AACGCUAC |
| 95 | UAGCGUUUC GUUGCUGU | ACAGCAAC CUGAUGA X GAA AAACGCUA |
| 98 | CGUUUCGUU GCUGUCAA | UUGACAGC CUGAUGA X GAA ACGAAACG |
| 104 | GUUGCUGUC AAAGCUCC | GGAGCUUU CUGAUGA X GAA ACAGCAAC |
| 111 | UCAAAGCUC CGCUCUCG | CGAGAGCG CUGAUGA X GAA AGCUUUGA |
| | GCUCCGCUC UCGAGCGG | CCGCUCGA CUGAUGA X GAA AGCGGAGC |
| 116 | UCCGCUCUC GAGCGGUC | GACCGCUC CUGAUGA X GAA AGAGCGGA |
| 118 | CGAGCGGUC CAACAGUC | GACUGUUG CUGAUGA X GAA ACCGCUCG |
| 126 | CCAACAGUC AAAUGUUA | UAACAUUU CUGAUGA X GAA ACUGUUGG |
| 134 | UCAAAUGUU AGCAAUUC | GAAUUGCU CUGAUGA X GAA ACAUUUGA |
| 141 | CAAAUGUUA GCAAUUCU | AGAAUUGC CUGAUGA X GAA AACAUUUG |
| 142 | UUAGCAAUU CUGUGCGC | GCGCACAG CUGAUGA X GAA AUUGCUAA |
| 148 | UAGCAAUUC UGUGCGCU | AGCGCACA CUGAUGA X GAA AAUUGCUA |
| 149 | | UGGACUUG CUGAUGA X GAA AGCCAGCG |
| 162 | GCUGGCUU CAAGUCCAA | UUGGACUU CUGAUGA X GAA AAGCCAGC |
| 163 | | GAGGUUG CUGAUGA X GAA ACUUGAAG |
| 168 | CUUCAAGUC CAAACCUC | GACCGGAA CUGAUGA X GAA AGGUUUGG |
| 176 | CCAAACCUC UUCCGGUC | AAGACCGG CUGAUGA X GAA AGAGGUUU |
| 178 | AAACCUCUU CCGGUCUU | CAAGACCG CUGAUGA X GAA AAGAGGUU |
| 179 | AACCUCUUC CGGUCUUG | CAGAUCAA CUGAUGA X GAA ACCGGAAG |
| 184 | CUUCCGGUC UUGAUCUG | CAGAUCAA CUGAUGA X GAA ACCGGAAG |
| 186 | UCCGGUCUU GAUCUGCG | AGAACGCA CUGAUGA X GAA AUCAAGAC |
| 190 - | GUCUUGAUC UGCGUUCU | CAGCUCAG CUGAUGA X GAA ACCAGAU |
| 196 | AUCUGCGUU CUGAGCUG | |
| 197 | UCUGCGUUC UGAGCUGG | CCAGCUCA CUGAUGA X GAA AACGCAGA |
| 207 | GAGCUGGUA CAAGAAUU | AAUUCUUG CUGAUGA X GAA ACCAGCUC |

Table V

| Nt. | Substrate | Ribozyme |
|------------|-----------------------|----------------------------------|
| Position | | |
| 215 | ACAAGAAUU GAUUCCUG | CAGGAAUC CUGAUGA X GAA AUUCUUGU |
| 219 | GAAUUGAUU CCUGAACA | UGUUCAGG CUGAUGA X GAA AUCAAUUC |
| 220 | AAUUGAUUC CUGAACAA | UUGUUCAG CUGAUGA X GAA AAUCAAUU |
| 235 | AACAGGAUC GCCUGAAA | UUUCAGGC CUGAUGA X GAA AUCCUGUU |
| 249 | AAAAAGAUC AAGUCAGA | UCUGACUU CUGAUGA X GAA AUCUUUUU |
| 254 | GAUCAAGUC AGAUAUGA | UCAUAUCU CUGAUGA X GAA ACUUGAUC |
| 259 | AGUCAGAUA UGAAAGGU | ACCUUUCA CUGAUGA X GAA AUCUGACU |
| 268 | UGAAAGGUU CAAUUGGG | CCCAAUUG CUGAUGA X GAA ACCUUUCA |
| | GAAAGGUUC AAUUGGGA | UCCCAAUU CUGAUGA X GAA AACCUUUC |
| 269 273 | GGUUCAAUU GGGAACAU | AUGUUCCC CUGAUGA X GAA AUUGAACC |
| | GGGAACAUC ACAGUUGA | UCAACUGU CUGAUGA X GAA AUGUUCCC |
| 282 | AUCACAGUU GAUAUGGU | ACCAUAUC CUGAUGA X GAA ACUGUGAU |
| 288 | | AAGAACCA CUGAUGA X GAA AUCAACUG |
| 292 | CAGUUGAUA UGGUUCUU | CCACCAAG CUGAUGA X GAA ACCAUAUC |
| 297 | GAUAUGGUU CUUGGUGG | UCCACCAA CUGAUGA X GAA AACCAUAU |
| 298 | AUAUGGUUC UUGGUGGA | AUUCCACC CUGAUGA X GAA AACCAUAU |
| 300 | AUGGUUCUU GGUGGAAU | UCCACAGU CUGAUGA X GAA AUCCUGUC |
| 326 | GACAGGAUU ACUGUGGA | UUCCACAGU CUGAUGA X GAA AAUCCUGU |
| 327 | ACAGGAUUA CUGUGGAA | UUCCACAG CUGAUGA X GAA AACCCUGU |
| 340 | GGAAACCUC AUUACCUU | AAGGUAAU CUGAUGA X GAA AGGUUUCC |
| 343 | AACCUCAUU ACCUUGAC | GUCAAGGU CUGAUGA X GAA AUGAGGUU |
| 344 | ACCUCAUUA CCUUGACC | 0000:2:00 |
| 348 | CAUUACCUU GACCCUGA | UCAGGGUC CUGAUGA X GAA AGGUAAUG |
| 366 | GAGGGAAUU CGCUUCCG | CGGAAGCG CUGAUGA X GAA AUUCCCUC |
| 367 | AGGGAAUUC GCUUCCGG | CCGGAAGC CUGAUGA X GAA AAUUCCCU |
| 371 | AAUUCGCUU CCGGGGGU | ACCCCCGG CUGAUGA X GAA AGCGAAUU |
| 372 | AUUCGCUUC CGGGGGUU | AACCCCCG CUGAUGA X GAA AAGCGAAU |
| 380 | CCGGGGGUU GUCUAUAC | GUAUAGAC CUGAUGA X GAA ACCCCCGG |
| 383 | GGGGUUGUC UAUACCUG | CAGGUAUA CUGAUGA X GAA ACAACCCC |
| ·385 | GGUUGUCUA UACCUGAA | UUCAGGUA CUGAUGA X GAA AGACAACC |
| 387 | UUGUCUAUA CCUGAAUG | CAUUCAGG CUGAUGA X GAA AUAGACAA |
| 405 | CAAAAGGUA UUACCUGC | GCAGGUAA CUGAUGA X GAA ACCUUUUG |
| 407 | AAAGGUAUU ACCUGCAG | CUGCAGGU CUGAUGA X GAA AUACCUUU |
| 408 | AAGGUAUUA CCUGCAGC | GCUGCAGG CUGAUGA X GAA AAUACCUU |
| 437 | UGAGCCCUU GCCUGAAG | CUUCAGGC CUGAUGA X GAA AGGGCUCA |
| 448 | CUGAAGGUC UUCUCUGG | CCAGAGAA CUGAUGA X GAA ACCUUCAG |
| 450 | GAAGGUCUU CUCUGGCU | AGCCAGAG CUGAUGA X GAA AGACCUUC |
| 451 | AAGGUCUUC UCUGGCUU | AAGCCAGA CUGAUGA X GAA AAGACCUU |
| 453 | GGUCUUCUC UGGCUUCU | AGAAGCCA CUGAUGA X GAA AGAAGACC |
| 459 | CUCUGGCUU CUUUUAAC | GUUAAAAG CUGAUGA X GAA AGCCAGAG |
| 460 | UCUGGCUUC UUUUAACA | UGUUAAAA CUGAUGA X GAA AAGCCAGA |
| 462 | UGGCUUCUU UUAACAGG | CCUGUUAA CUGAUGA X GAA AGAAGCCA |
| 463 | GGCUUCUUU UAACAGGA | UCCUGUUA CUGAUGA X GAA AAGAAGCC |
| 464 | GCUUCUUUU AACAGGAA | UUCCUGUU CUGAUGA X GAA AAAGAAGC |
| 465 | CUUCUUUUA ACAGGAAA | UUUCCUGU CUGAUGA X GAA AAAAGAAG |
| 482 | GGUGCCAUC AAAAGAGC | GCUCUUUU CUGAUGA X GAA AUGGCACC |
| 499 | AAGUGAAUU CAAUUGUC | GACAAUUG CUGAUGA X GAA AUUCACUU |
| 500 | AGUGAAUUC AAUUGUCU | AGACAAUU CUGAUGA X GAA AAUUCACU |
| 504 | AAUUCAAUU GUCUCAGG | CCUGAGAC CUGAUGA X GAA AUUGAAUU |
| 507 | UCAAUUGUC UCAGGAAU | AUUCCUGA CUGAUGA X GAA ACAAUUGA |
| | AAUUGUCUC AGGAAUUG | CAAUUCCU CUGAUGA X GAA AGACAAUU |
| 509 | UCAGGAAUU GCAGAGUC | GACUCUGC CUGAUGA X GAA AUUCCUGA |
| 516 | UGCAGAGUC GGGCAUCA | UGAUGCC CUGAUGA X GAA ACUCUGCA |
| 524 | UCGGGCAUC AUAUCCCU | AGGGAUAU CUGAUGA X GAA AUGCCCGA |
| 531 | COCATICATA DECEDICATI | AUCAGGGA CUGAUGA X GAA AUGAUGCC |
| 534 | GGCAUCAUA UCCCUGAU | UGAUCAGG CUGAUGA X GAA AUAUGAUG |
| 536 | CAUCAUAUC CCUGAUCA | UACAUGAU CUGAUGA X GAA AUCAGGGA |
| 543 | UCCCUGAUC AUCAUGUA | GUAUACAU CUGAUGA X GAA AUGAUCAG |
| 546 | CUGAUCAUC AUGUAUAC | GUMUNCHU CUGNUGN A GNA AUGNOCHG |

Table V

| Nt. | Substrate | Ribozyme |
|----------|--------------------|--|
| Position | | |
| 551 | CAUCAUGUA UACAACUA | UAGUUGUA CUGAUGA X GAA ACAUGAUG |
| 553 | UCAUGUAUA CAACUAUU | AAUAGUUG CUGAUGA X GAA AUACAUGA |
| 559 | AUACAACUA UUGAUGCC | GGCAUCAA CUGAUGA X GAA AGUUGUAU |
| 561 | ACAACUAUU GAUGCCUU | AAGGCAUC CUGAUGA X GAA AUAGUUGU |
| 569 | UGAUGCCUU ACCAGUCA | UGACUGGU CUGAUGA X GAA AGGCAUCA |
| | GAUGCCUUA CCAGUCAC | GUGACUGG CUGAUGA X GAA AAGGCAUC |
| 570 | UUACCAGUC ACAGCUCA | UGAGCUGU CUGAUGA X GAA ACUGGUAA |
| 576 | UCACAGCUC AUCCAAUG | CAUUGGAU CUGAUGA X GAA AGCUGUGA |
| 583 | CAGCUCAUC CAAUGACC | GGUCAUUG CUGAUGA X GAA AUGAGCUG |
| 586 | 0.10000 | CAGUAGCA CUGAUGA X GAA ACUGGGUC |
| 599 | 0 | CCAGUAGC CUGAUGA X GAA AACUGGGU |
| 600 | | GACUCCAG CUGAUGA X GAA AGCAAACU |
| 604 | AGUUUGCUA CUGGAGUC | AGAGCCAU CUGAUGA X GAA ACUCCAGU |
| 612 | ACUGGAGUC AUGGCUCU | AACCUGAA CUGAUGA X GAA AGCCAUGA |
| 619 | UCAUGGCUC UUCAGGUU | |
| 621 | AUGGCUCUU CAGGUUCA | COMMODULE CONTRACTOR |
| 622 | UGGCUCUUC AGGUUCAA | 000.2.200 |
| 627 | CUUCAGGUU CAAAGUGA | 00.10000 |
| 628 | UUCAGGUUC AAAGUGAA | OCCABODO COSITORIO |
| 638 | AAGUGAAUU UCAAAAGG | |
| 639 | AGUGAAUUU CAAAAGGC | 0000000 |
| 640 | GUGAAUUUC AAAAGGCA | 0000000 000110111 |
| 650 | AAAGGCAUA CGAGAAAG | CUUUCUCG CUGAUGA X GAA AUGCCUUU |
| 663 | AAAGGGAUU CACAAAUC | GAUUUGUG CUGAUGA X GAA AUCCCUUU |
| 664 | AAGGGAUUC ACAAAUCA | UGAUUUGU CUGAUGA X GAA AAUCCCUU |
| 671 | UCACAAAUC AAAGUAUU | AAUACUUU CUGAUGA X GAA AUUUGUGA |
| 677 | AUCAAAGUA UUGGGAAC | GUUCCCAA CUGAUGA X GAA ACUUUGAU |
| 679 | CAAAGUAUU GGGAACCA | UGGUUCCC CUGAUGA X GAA AUACUUUG |
| 692 | ACCAACAUA UGAGGAUU | AAUCCUCA CUGAUGA X GAA AUGUUGGU |
| 700 | AUGAGGAUU CCAUGAAU | AUUCAUGG CUGAUGA X GAA AUCCUCAU |
| 701 | UGAGGAUUC CAUGAAUC | GAUUCAUG CUGAUGA X GAA AAUCCUCA |
| 709 | CCAUGAAUC UGAUUGCU | AGCAAUCA CUGAUGA X GAA AUUCAUGG |
| 714 | AAUCUGAUU GCUCAAGU | ACUUGAGC CUGAUGA X GAA AUCAGAUU |
| 718 | UGAUUGCUC AAGUUCCA | UGGAACUU CUGAUGA X GAA AGCAAUCA |
| 723 | GCUCAAGUU CCACUUGU | ACAAGUGG CUGAUGA X GAA ACUUGAGC |
| 724 | CUCAAGUUC CACUUGUU | AACAAGUG CUGAUGA X GAA AACUUGAG |
| 729 | GUUCCACUU GUUGCUGC | GCAGCAAC CUGAUGA X GAA AGUGGAAC |
| 732 | CCACUUGUU GCUGCUUA | UAAGCAGC CUGAUGA X GAA ACAAGUGG |
| 739 | UUGCUGCUU AUGUUUAU | AUAAACAU CUGAUGA X GAA AGCAGCAA |
| 740 | UGCUGCUUA UGUUUAUC | GAUAAACA CUGAUGA X GAA AAGCAGCA |
| 744 | GCUUAUGUU UAUCGCAG | CUGCGAUA CUGAUGA X GAA ACAUAAGC |
| 745 | CUUAUGUUU AUCGCAGG | CCUGCGAU CUGAUGA X GAA AACAUAAG |
| 746 | UUAUGUUUA UCGCAGGA | UCCUGCGA CUGAUGA X GAA AAACAUAA |
| 748 | AUGUUUAUC GCAGGAUG | CAUCCUGC CUGAUGA X GAA AUAAACAU |
| 758 | CAGGAUGUA CAAGAAUG | CAUUCUUG CUGAUGA X GAA ACAUCCUG |
| | GUGACACUA UACCUAAG | CUUAGGUA CUGAUGA X GAA AGUGUCAC |
| 775 | GACACUAUA CCUAAGGA | UCCUUAGG CUGAUGA X GAA AUAGUGUC |
| | CUAUACCUA AGGAUGAA | UUCAUCCU CUGAUGA X GAA AGGUAUAG |
| 781 | GGAUGAAUC CCUGGAUU | AAUCCAGG CUGAUGA X GAA AUUCAUCC |
| 791 | CCCUGGAUU AUGGUGCA | UGCACCAU CUGAUGA X GAA AUCCAGGG |
| 799 | CCUGGAUUA UGGUGCAA | UUGCACCA CUGAUGA X GAA AAUCCAGG |
| 800 | CUUGGAUUA UGGUGCAA | GUGAGCA CUGAUGA X GAA AUUUGCAC |
| 811 | GUGCAAAUU UUGCUCAC | UGUGAGCA CUGAUGA X GAA AAUUUGCA |
| 812 | UGCAAAUUU UGCUCACA | AUGUGAGC CUGAUGA X GAA AAAUUUGC |
| 813 | GCAAAUUUU GCUCACAU | AAGCAUGU CUGAUGA X GAA AGCAAAAU |
| 817 | AUUUUGCUC ACAUGCUU | CUGAAACC CUGAUGA X GAA AGCAUGUG |
| 825 | CACAUGCUU GGUUUCAG | |
| 829 | UGCUUGGUU UCAGUAGC | GCUACUGA CUGAUGA X GAA ACCAAGCA AGCUACUG CUGAUGA X GAA AACCAAGC |
| | GCUUGGUUU CAGUAGCU | |

Table V

| Nt. | Substrate | Ribozyme |
|----------|--------------------|---------------------------------|
| Position | | - CDD DDDCCDDC |
| 831 | CUUGGUUUC AGUAGCUC | GAGCUACU CUGAUGA X GAA AAACCAAG |
| 835 | GUUUCAGUA GCUCUGAA | UUCAGAGC CUGAUGA X GAA ACUGAAAC |
| 839 | CAGUAGCUC UGAAAUGC | GCAUUUCA CUGAUGA X GAA AGCUACUG |
| 855 | CAUGAACUU CUUAUGAG | CUCAUAAG CUGAUGA X GAA AGUUCAUG |
| 856 | AUGAACUUC UUAUGAGG | CCUCAUAA CUGAUGA X GAA AAGUUCAU |
| 858 | GAACUUCUU AUGAGGCU | AGCCUCAU CUGAUGA X GAA AGAAGUUC |
| 859 | AACUUCUUA UGAGGCUC | GAGCCUCA CUGAUGA X GAA AAGAAGUU |
| 867 | AUGAGGCUC UAUGUAAC | GUUACAUA CUGAUGA X GAA AGCCUCAU |
| 869 | GAGGCUCUA UGUAACAA | UUGUUACA CUGAUGA X GAA AGAGCCUC |
| | CUCUAUGUA ACAAUACA | UGUAUUGU CUGAUGA X GAA ACAUAGAG |
| 873 | GUAACAAUA CACAGUGA | UCACUGUG CUGAUGA X GAA AUUGUUAC |
| 879 | ACAGUGAUC AUGAAGGU | ACCUUCAU CUGAUGA X GAA AUCACUGU |
| 889 | | ACUGACAU CUGAUGA X GAA ACCACCUU |
| 901 | AAGGUGGUA AUGUCAGU | UGAGCACU CUGAUGA X GAA ACAUUACC |
| 906 | GGUAAUGUC AGUGCUCA | 00.100.100 |
| 913 | UCAGUGCUC ACACCGGU | |
| 922 | ACACCGGUC ACUUGGUU | 1210012100 |
| 926 | CGGUCACUU GGUUGCUA | UAGCAACC CUGAUGA X GAA AGUGACCG |
| 930 | CACUUGGUU GCUAGUGC | GCACUAGC CUGAUGA X GAA ACCAAGUG |
| 934 | UGGUUGCUA GUGCUUUG | CAAAGCAC CUGAUGA X GAA AGCAACCA |
| 940 | CUAGUGCUU UGUCUGAU | AUCAGACA CUGAUGA X GAA AGCACUAG |
| 941 | UAGUGCUUU GUCUGAUC | GAUCAGAC CUGAUGA X GAA AAGCACUA |
| 944 | UGCUUUGUC UGAUCCUU | AAGGAUCA CUGAUGA X GAA ACAAAGCA |
| 949 | UGUCUGAUC CUUACCUC | GAGGUAAG CUGAUGA X GAA AUCAGACA |
| | CUGAUCCUU ACCUCUCC | GGAGAGGU CUGAUGA X GAA AGGAUCAG |
| 952 | UGAUCCUUA CCUCUCCU | AGGAGAGG CUGAUGA X GAA AAGGAUCA |
| 953 | CCUUACCUC UCCUUUGC | GCAAAGGA CUGAUGA X GAA AGGUAAGG |
| 957 | | CAGCAAAG CUGAUGA X GAA AGAGGUAA |
| 959 | | CAGCAGCA CUGAUGA X GAA AGGAGAGG |
| 962 | CCUCUCCUU UGCUGCUG | 0001.01 |
| 963 | CUCUCCUUU GCUGCUGC | 0011001100 0001110011 |
| 973 | CUGCUGCUU UGAAUGGU | riceriocer: courte |
| 974 | UGCUGCUUU GAAUGGUU | |
| 982 | UGAAUGGUU UAGCCGGA | UCCGGCUA CUGAUGA X GAA ACCAUUCA |
| 983 | GAAUGGUUU AGCCGGAC | GUCCGGCU CUGAUGA X GAA AACCAUUC |
| 984 | AAUGGUUUA GCCGGACC | GGUCCGGC CUGAUGA X GAA AAACCAUU |
| 996 | GGACCACUU CAUGGUUU | AAACCAUG CUGAUGA X GAA AGUGGUCC |
| 997 | GACCACUUC AUGGUUUA | UAAACCAU CUGAUGA X GAA AAGUGGUC |
| 600 | UUCAUGGUU UAGCCAAU | AUUGGCUA CUGAUGA X GAA ACCAUGAA |
| 1004 | UCAUGGUUU AGCCAAUC | GAUUGGCU CUGAUGA X GAA AACCAUGA |
| 1005 | CAUGGUUUA GCCAAUCA | UGAUUGGC CUGAUGA X GAA AAACCAUG |
| 1012 | UAGCCAAUC AGGAAGUU | AACUUCCU CUGAUGA X GAA AUUGGCUA |
| 1020 | CAGGAAGUU UUGCUAUG | CAUAGCAA CUGAUGA X GAA ACUUCCUG |
| | AGGAAGUUU UGCUAUGG | CCAUAGCA CUGAUGA X GAA AACUUCCU |
| 1021 | GGAAGUUUU GCUAUGGA | UCCAUAGC CUGAUGA X GAA AAACUUCC |
| | GUUUUGCUA UGGAUAAA | UUUAUCCA CUGAUGA X GAA AGCAAAAC |
| 1026 | CUAUGGAUA AAAUCUGU | ACAGAUUU CUGAUGA X GAA AUCCAUAG |
| 1032 | CUAUGGAUA AAAUCUGU | CUACAACA CUGAUGA X GAA AUUUUAUC |
| 1037 | GAUAAAAUC UGUUGUAG | UCUUCUAC CUGAUGA X GAA ACAGAUUU |
| 1041 | AAAUCUGUU GUAGAAGA | 00000000 |
| 1044 | UCUGUUGUA GAAGAAUG | Criococc coefficient |
| 1065 | GAGAACAUU UCCAAAGA | |
| 1066 | AGAACAUUU CCAAAGAG | |
| 1067 | GAACAUUUC CAAAGAGC | GCUCUUUG CUGAUGA X GAA AAAUGUUC |
| 1079 | AGAGCAGUU GAAAGACU | AGUCUUUC CUGAUGA X GAA ACUGCUCU |
| 1088 | GAAAGACUA UGUUUGGA | UCCAAACA CUGAUGA X GAA AGUCUUUC |
| 1092 | GACUAUGUU UGGAAAAC | GUUUUCCA CUGAUGA X GAA ACAUAGUC |
| 1093 | ACUAUGUUU GGAAAACA | UGUUUUCC CUGAUGA X GAA AACAUAGU |
| 1103 | GAAAACAUU GAACAGUG | CACUGUUC CUGAUGA X GAA AUGUUUUC |
| 1100 | GGCAAGGUU GUCCCUGG | CCAGGGAC CUGAUGA X GAA ACCUUGCC |

Table V

| | | Diversion |
|------------------|--------------------|---|
| Nt. | Substrate | Ribozyme |
| Position 1122 | AAGGUUGUC CCUGGUUU | AAACCAGG CUGAUGA X GAA ACAACCUU |
| 1122 | UCCCUGGUU UUGGACAU | AUGUCCAA CUGAUGA X GAA ACCAGGGA |
| 1130 | CCCUGGUUU UGGACAUG | CAUGUCCA CUGAUGA X GAA AACCAGGG |
| 1131 | CCUGGUUUU GGACAUGG | CCAUGUCC CUGAUGA X GAA AAACCAGG |
| 1143 | CAUGGAGUU CUGCGAAA | UUUCGCAG CUGAUGA X GAA ACUCCAUG |
| 1143 | AUGGAGUUC UGCGAAAG | CUUUCGCA CUGAUGA X GAA AACUCCAU |
| 1158 | AAGACUGUA CCAAGAUA | UAUCUUGG CUGAUGA X GAA ACAGUCUU |
| 1166 | ACCAAGAUA UACAUGCC | GGCAUGUA CUGAUGA X GAA AUCUUGGU |
| 1168 | CAAGAUAUA CAUGCCAG | CUGGCAUG CUGAUGA X GAA AUAUCUUG |
| 1184 | GAGAGAGUU CGCUAUGA | UCAUAGOG CUGAUGA X GAA ACUCUCUC |
| 1185 | AGAGAGUUC GCUAUGAA | UUCAUAGC CUGAUGA X GAA AACUCUCU |
| 1189 | AGUUCGCUA UGAAGCAU | AUGCUUCA CUGAUGA X GAA AGCGAACU |
| 1198 | UGAAGCAUU UGCCUGAA | UUCAGGCA CUGAUGA X GAA AUGCUUCA |
| 1199 | GAAGCAUUU GCCUGAAG | CUUCAGGC CUGAUGA X GAA AAUGCUUC |
| 1210 | CUGAAGAUC CACUGUUU | AAACAGUG CUGAUGA X GAA AUCUUCAG |
| | UCCACUGUU UCAACUGG | CCAGUUGA CUGAUGA X GAA ACAGUGGA |
| 1217 | CCACUGUUU CAACUGGU | ACCAGUUG CUGAUGA X GAA AACAGUGG |
| 1218 | CACUGUUUC AACUGGUU | AACCAGUU CUGAUGA X GAA AAACAGUG |
| 1227 | CAACUGGUU UCAAAACU | AGUUUUGA CUGAUGA X GAA ACCAGUUG |
| 1228 | AACUGGUUU CAAAACUC | GAGUUUUG CUGAUGA X GAA AACCAGUU |
| 1229 | ACUGGUUUC AAAACUCU | AGAGUUUU CUGAUGA X GAA AAACCAGU |
| 1236 | UCAAAACUC UACGAAGU | ACUUCGUA CUGAUGA X GAA AGUUUUGA |
| 1238 | AAAACUCUA CGAAGUUU | AAACUUCG CUGAUGA X GAA AGAGUUUU |
| 1245 | UACGAAGUU UUCCUCCU | AGGAGGAA CUGAUGA X GAA ACUUCGUA |
| 1245 | ACGAAGUUU UCCUCCUG | CAGGAGGA CUGAUGA X GAA AACUUCGU |
| 1247 | CGAAGUUUU CCUCCUGU | ACAGGAGG CUGAUGA X GAA AAACUUCG |
| 1248 | GAAGUUUUC CUCCUGUU | AACAGGAG CUGAUGA X GAA AAAACUUC |
| 1251 | GUUUUCCUC CUGUUCUU | AAGAACAG CUGAUGA X GAA AGGAAAAC |
| 1256 | CCUCCUGUU CUUACAGA | UCUGUAAG CUGAUGA X GAA ACAGGAGG |
| 1257 | CUCCUGUUC UUACAGAA | UUCUGUAA CUGAUGA X GAA AACAGGAG |
| 1259 | CCUGUUCUU ACAGAACU | AGUUCUGU CUGAUGA X GAA AGAACAGG |
| 1260 | CUGUUCUUA CAGAACUU | AAGUUCUG CUGAUGA X GAA AAGAACAG |
| 1268 | ACAGAACUU GGCAAAGU | ACUUUGCC CUGAUGA X GAA AGUUCUGU |
| 1277 | GGCAAAGUU AAAACCUU | AAGGUUUU CUGAUGA X GAA ACUUUGCC |
| 1278 | GCAAAGUUA AAACCUUG | CAAGGUUU CUGAUGA X GAA AACUUUGC |
| 1285 | UAAAACCUU GGCCAAAU | AUUUGGCC CUGAUGA X GAA AGGUUUUA |
| 1296 | CCAAAUGUU GAUGCCCA | UGGGCAUC CUGAUGA X GAA ACAUUUGG |
| 1316 | UGGUGUGUU GUUGAACU | AGUUCAAC CUGAUGA X GAA ACACACCA |
| 1319 | UGUGUUGUU GAACUAUU | AAUAGUUC CUGAUGA X GAA ACAACACA |
| 1325 | GUUGAACUA UUAUGGUU | AACCAUAA CUGAUGA X GAA AGUUCAAC |
| 1327 | UGAACUAUU AUGGUUUA | UAAACCAU CUGAUGA X GAA AUAGUUCA |
| 1328 | GAACUAUUA UGGUUUAA | UUAAACCA CUGAUGA X GAA AAUAGUUC |
| 1333 | AUUAUGGUU UAACUGAA | UUCAGUUA CUGAUGA X GAA ACCAUAAU |
| 1334 | UUAUGGUUU AACUGAAG | CUUCAGUU CUGAUGA X GAA AACCAUAA |
| 1335 | UAUGGUUUA ACUGAAGC | GCUUCAGU CUGAUGA X GAA AAACCAUA |
| 1349 | AGCAAGAUA UUAUACGG | CCGUAUAA CUGAUGA X GAA AUCUUGCU |
| 1351 | CAAGAUAUU AUACGGUC | GACCGUAU CUGAUGA X GAA AUAUCUUG |
| 1352 | AAGAUAUUA UACGGUCC | GGACCGUA CUGAUGA X GAA AAUAUCUU |
| 1354 | GAUAUUAUA CGGUCCUC | GAGGACCG CUGAUGA X GAA AUAAUAUC |
| 1359 | UAUACGGUC CUCUUUGG | CCAAAGAG CUGAUGA X GAA ACCGUAUA |
| 1362 | ACGGUCCUC UUUGGCGU | ACGCCAAA CUGAUGA X GAA AGGACCGU |
| 1364 | GGUCCUCUU UGGCGUAU | AUACGCCA CUGAUGA X GAA AGAGGACC |
| 1365 | GUCCUCUUU GGCGUAUC | GAUACGCC CUGAUGA X GAA AAGAGGAC |
| 1371 | UUUGGCGUA UCAAGAGC | GCUCUUGA CUGAUGA X GAA ACGCCAAA |
| 1373 | UGGCGUAUC AAGAGCUC | GAGCUCUU CUGAUGA X GAA AUACGCCA AAUGCCAA CUGAUGA X GAA AGCUCJUG |
| 1381 | CAAGAGCUC UUGGCAUU | |
| 1383 | AGAGCUCUU GGCAUUUG | CAAAUGCC CUGAUGA X GAA AGAGCUCU |

Table V

| Nt. | Substrate | Ribozyme | |
|--------------|--------------------|--|--|
| Position | | THE PARTY OF THE P | |
| 1389 | CUUGGCAUU UGCUCUCA | UGAGAGCA CUGAUGA X GAA AUGCCAAG CUGAGAGC CUGAUGA X GAA AAUGCCAA | |
| 1390 | UUGGCAUUU GCUCUCAG | 00010101 | |
| 1394 | CAUUUGCUC UCAGCUAA | 001100001 | |
| 1396 | UUUGCUCUC AGCUAAUU | | |
| 1401 | UCUCAGCUA AUUUGGGA | UCCCAAAU CUGAUGA X GAA AGCUGAGA | |
| 1404 | CAGCUAAUU UGGGACCG | CGGUCCCA CUGAUGA X GAA AUUAGCUG UCGGUCCC CUGAUGA X GAA AAUUAGCU | |
| 1405 | AGCUAAUUU GGGACCGA | | |
| 1417 | ACCGAGCUC UUGGAUUG | 0.1.000 | |
| 1419 | CGAGCUCUU GGAUUGCC | GGCAAGCC CCCIICCII II CI | |
| 1424 | UCUUGGAUU GCCGCUAG | | |
| 1431 | UUGCCGCUA GAGAGGCC | | |
| 1449 | AAGAGUGUC ACAAUGGA | | |
| 1464 | GAGUGGCUU GAGAACCA | | |
| 1491 | GCAUGAAUU GUUUGAAA | | |
| 1494 | UGAAUUGUU UGAAAUCU | AGAUUUCA CUGAUGA X GAA ACAAUUCA | |
| 1495 | GAAUUGUUU GAAAUCUC | GAGAUUUC CUGAUGA X GAA AACAAUUC | |
| 1501 | UUUGAAAUC UCGCGAGC | GCUCGCGA CUGAUGA X GAA AUUUCAAA | |
| 1503 | UGAAAUCUC GCGAGCAU | AUGCUCGC CUGAUGA X GAA AGAUUUCA | |
| 1512 | GCGAGCAUA AAACACAA | UUGUGUUU CUGAUGA X GAA AUGCUCGC | |
| 1524 | CACAAUGUA UAAUCUCU | AGAGAUUA CUGAUGA X GAA ACAUUGUG | |
| 1526 | CAAUGUAUA AUCUCUAU | AUAGAGAU CUGAUGA X GAA AUACAUUG | |
| 1529 | UGUAUAAUC UCUAUGAA | UUCAUAGA CUGAUGA X GAA AUUAUACA | |
| 1531 | UAUAAUCUC UAUGAAUA | UAUUCAUA CUGAUGA X GAA AGAUUAUA | |
| 1533 | UAAUCUCUA UGAAUAAU | AUUAUUCA CUGAUGA X GAA AGAGAUUA | |
| 1539 | CUAUGAAUA AUUGCUUG | CAAGCAAU CUGAUGA X GAA AUUCAUAG | |
| 1542 | UGAAUAAUU GCUUGACA | UGUCAAGC CUGAUGA X GAA AUUAUUCA | |
| 1546 | UAAUUGCUU GACAAAGC | GCUUUGUC CUGAUGA X GAA AGCAAUUA | |
| 1558 | AAAGCACUC CUUUCUUG | CAAGAAAG CUGAUGA X GAA AGUGCUUU | |
| 1561 | GCACUCCUU UCUUGGGG | CCCCAAGA CUGAUGA X GAA AGGAGUGC | |
| 1562 | CACUCCUUU CUUGGGGG | CCCCCAAG CUGAUGA X GAA AAGGAGUG | |
| 1563 | ACUCCUUUC UUGGGGGA | UCCCCCAA CUGAUGA X GAA AAAGGAGU | |
| 1565 | UCCUUUCUU GGGGGACA | UGUCCCCC CUGAUGA X GAA AGAAAGGA | |
| 1578 | GACAAGAUA GGUCGGCC | GGCCGACC CUGAUGA X GAA AUCUUGUC | |
| 1582 | AGAUAGGUC GGCCCUUC | GAAGGGCC CUGAUGA X GAA ACCUAUCU | |
| 1589 | UCGGCCCUU CAAUGGGU | ACCCAUUG CUGAUGA X GAA AGGGCCGA | |
| 1590 | CGGCCCUUC AAUGGGUU | AACCCAUU CUGAUGA X GAA AAGGGCCG | |
| 1598 | CAAUGGGUU AACGAACU | AGUUCGUU CUGAUGA X GAA ACCCAUUG | |
| 1599 | AAUGGGUUA ACGAACUU | AAGUUCGU CUGAUGA X GAA AACCCAUU | |
| 1607 | AACGAACUU CAGUUCAA | UUGAACUG CUGAUGA X GAA AGUUCGUU | |
| 1608 | ACGAACUUC AGUUCAAA | UUUGAACU CUGAUGA X GAA AAGUUCGU | |
| 1612 | ACUUCAGUU CAAACUUC | GAAGUUUG CUGAUGA X GAA ACUGAAGU | |
| 1613 | CUUCAGUUC AAACUUCA | UGAAGUUU CUGAUGA X GAA AACUGAAG | |
| 1619 | UUCAAACUU CACUGAAU | AUUCAGUG CUGAUGA X GAA AGUUUGAA | |
| 1620 | UCAAACUUC ACUGAAUU | AAUUCAGU CUGAUGA X GAA AAGUUUGA | |
| | CACUGAAUU UGUGUGAA | UUCACACA CUGAUGA X GAA AUUCAGUG | |
| 1628 1629 | ACUGAAUUU GUGUGAAU | AUUCACAC CUGAUGA X GAA AAUUCAGU | |
| | GUGUGAAUU GUAUGGUU | AACCAUAC CUGAUGA X GAA AUUCACAC | |
| 1638 | UGAAUUGUA UGGUUUCU | AGAAACCA CUGAUGA X GAA ACAAUUCA | |
| 1641 | UGUAUGGUU UCUCGAGA | UCUCGAGA CUGAUGA X GAA ACCAUACA | |
| | GUAUGGUUU CUCGAGAC | GUCUCGAG CUGAUGA X GAA AACCAUAC | |
| 1647 | UAUGGUUUC UCGAGACU | AGUCUCGA CUGAUGA X GAA AAACCAUA | |
| 1648 | UGGUUUCUC GAGACUUG | CAAGUCUC CUGAUGA X GAA AGAAACCA | |
| 1650 | UCGAGACUU GUCCUGAA | UUCAGGAC CUGAUGA X GAA AGUCUCGA | |
| 1657 | AGACUUGUC CUGAAUUU | AAAUUCAG CUGAUGA X GAA ACAAGUCU | |
| 1660 | AGACOUGUC COGAAOOO | AAGUUCAA CUGAUGA X GAA AUUCAGGA | |
| 1667 | UCCUGAAUU UUGAACUU | UAAGUUCA CUGAUGA X GAA AAUUCAGG | |
| 1668 | CCUGAAUUU UGAACUUA | CUAAGUUC CUGAUGA X GAA AAAUUCAG | |
| 1669 | CUGAAUUUU GAACUUAG | COMMODUC COGMOGN A GALL 122.500110 | |

Table V

| | 0.1 | Ribozyme | |
|----------|---------------------------------------|--|--|
| Nt. | Substrate | Ribozyme | |
| Position | UUUGAACUU AGUCUAGU | ACUAGACU CUGAUGA X GAA AGUUCAAA | |
| 1675 | | CACUAGAC CUGAUGA X GAA AAGUUCAA | |
| 1676 | V V V V V V V V V V V V V V V V V V V | AUCCACUA CUGAUGA X GAA ACUAAGUU | |
| 1679 | AACUUAGUC UAGUGGAU CUUAGUCUA GUGGAUUC | GAAUCCAC CUGAUGA X GAA AGACUAAG | |
| 1681 | | GAAAAAUG CUGAUGA X GAA AUCCACUA | |
| 1668 | 0.200 | AGAAAAAU CUGAUGA X GAA AAUCCACU | |
| 1689 | AGUGGAUUC AUUUUUCU | UGAAGAAA CUGAUGA X GAA AUGAAUCC | |
| 1692 | GGAUUCAUU UUUCUUCA | AUGAAGAA CUGAUGA X GAA AAUGAAUC | |
| 1693 | GAUUCAUUU UUCUUCAU | AAUGAAGA CUGAUGA X GAA AAAUGAAU | |
| 1694 | 1100011000 | GAAUGAAG CUGAUGA X GAA AAAAUGAA | |
| 1695 | UUCAUUUUU CUUCAUUC | 0.2.10 | |
| 1696 | UCAUUUUC UUCAUUCC | 0012100121 | |
| 1698 | AUUUUUCUU CAUUCCGA | COCCILIOS COCCI | |
| 1699 | UUUUUCUUC AUUCCGAA | 00001210 | |
| 1702 | UUCUUCAUU CCGAAUUC | GP2100CCC CCCITCCT | |
| 1703 | UCUUCAUUC CGAAUUCC | 00.2.0000 00 | |
| 1709 | UUCCGAAUU CCUCACAC | 000001100 000111011 | |
| 1710 | UCCGAAUUC CUCACACG | Chocochic Cochicon | |
| 1713 | GAAUUCCUC ACACGCUG | 0.100000 | |
| 1724 | ACGCUGAUC CAGCAUGU | | |
| 1733 | CAGCAUGUA AAAAUUAA | COLLIGORE COULT | |
| 1739 | GUAAAAAUU AAUAGGUC | 0.100 | |
| 1740 | UAAAAAUUA AUAGGUCA | UGACCUAU CUGAUGA X GAA AAUUUUUA | |
| 1743 | AAAUUAAUA GGUCAAUG | CAUUGACC CUGAUGA X GAA AUUAAUUU | |
| 1747 | UAAUAGGUC AAUGCUAU | AUAGCAUU CUGAUGA X GAA ACCUAUUA | |
| 1754 | UCAAUGCUA UUAAUCGC | GCGAUUAA CUGAUGA X GAA AGCAUUGA | |
| 1756 | AAUGCUAUU AAUCGCGU | ACGCGAUU CUGAUGA X GAA AUAGCAUU | |
| 1757 | AUGCUAUUA AUCGCGUU | AACGCGAU CUGAUGA X GAA AAUAGCAU | |
| 1760 | CUAUUAAUC GCGUUCUU | AAGAACGC CUGAUGA X GAA AUUAAUAG | |
| 1765 | AAUCGCGUU CUUGGUUG | CAACCAAG CUGAUGA X GAA ACGCGAUU | |
| 1766 | AUCGCGUUC UUGGUUGC | GCAACCAA CUGAUGA X GAA AACGCGAU | |
| 1768 | CGCGUUCUU GGUUGCCA | 000012100 | |
| 1772 | UUCUUGGUU GCCAUUAG | CUAAUGGC CUGAUGA X GAA ACCAAGAA | |
| 1778 | GUUGCCAUU AGACUUGU | ACAAGUCU CUGAUGA X GAA AUGGCAAC CACAAGUC CUGAUGA X GAA AAUGGCAA | |
| 1779 | UUGCCAUUA GACUUGUG | V. 10-11-11-11-11-11-11-11-11-11-11-11-11-1 | |
| 1784 | AUUAGACUU GUGAAUGA | 00:1000:10 | |
| 1795 | GAAUGACUU CCUUUGCU | AGCAAAGG CUGAUGA X GAA AGUCAUUC | |
| 1796 | AAUGACUUC CUUUGCUG | CAGCAAAG CUGAUGA X GAA AAGUCAUU | |
| 1799 | GACUUCCUU UGCUGGAA | 0000110011 | |
| 1800 | ACUUCCUUU GCUGGAAA | UUUCCAGC CUGAUGA X GAA AAGGAAGU | |
| 1811 | UGGAAAGUU AGUAAUCG | CGAUUACU CUGAUGA X GAA ACUUUCCA | |
| 1812 | GGAAAGUUA GUAAUCGG | | |
| 1815 | AAGUUAGUA AUCGGCUG | CAGCCGAU CUGAUGA X GAA ACUAACUU | |
| 1818 | UUAGUAAUC GGCUGAUU | AAUCAGCC CUGAUGA X GAA AUUACUAA AUUGCGUG CUGAUGA X GAA AUCAGCCG | |
| 1826 | CGGCUGAUU CACGCAAU | | |
| 1827 | GGCUGAUUC ACGCAAUA | 011000000 | |
| 1835 | CACGCAAUA AACUGCAA | 00001000 000110011 11 | |
| 1845 | ACUGCAAUU GUGUAGUU | AACUACAC CUGAUGA X GAA AUUGCAGU | |
| 1850 | AAUUGUGUA GUUUCUUA | UAAGAAAC CUGAUGA X GAA ACACAAUU | |
| 1853 | UGUGUAGUU UCUUAAAU | AUUUAAGA CUGAUGA X GAA ACUACACA | |
| 1854 | GUGUAGUUU CUUAAAUU | AAUUUAAG CUGAUGA X GAA AACUACAC | |
| 1855 | UGUAGUUUC UUAAAUUU | AAAUUUAA CUGAUGA X GAA AAACUACA | |
| 1857 | UAGUUUCUU AAAUUUGC | GCAAAUUU CUGAUGA X GAA AGAAACUA | |
| 1858 | AGUUUCUUA AAUUUGCU | AGCAAAUU CUGAUGA X GAA AAGAAACU | |
| 1862 | UCUUAAAUU UGCUAAUU | AAUUAGCA CUGAUGA X GAA AUUUAAGA | |
| 1863 | CUUAAAUUU GCUAAUUC | GAAUUAGC CUGAUGA X GAA AAUUUAAG | |
| 1867 | AAUUUGCUA AUUCUUAU | AUAAGAAU CUGAUGA X GAA AGCAAAUU | |
| 1870 | UUGCUAAUU CUUAUUUG | CAAAUAAG CUGAUGA X GAA AUUAGCAA | |

Table V

| Nt. | Substrate | Ribozyme | |
|----------|--------------------|--|--|
| Position | | THE PART OF THE PA | |
| 1871 | UGCUAAUUC UUAUUUGA | UCAAAUAA CUGAUGA X GAA AAUUAGCA | |
| 1873 | CUAAUUCUU AUUUGAUG | CAUCAAAU CUGAUGA X GAA AGAAUUAG | |
| 1874 | UAAUUCUUA UUUGAUGA | UCAUCAAA CUGAUGA X GAA AAGAAUUA | |
| 1876 | AUUCUUAUU UGAUGAUA | UAUCAUCA CUGAUGA X GAA AUAAGAAU | |
| | UUCUUAUUU GAUGAUAU | AUAUCAUC CUGAUGA X GAA AAUAAGAA | |
| 1877 | UUCUUAUUU GAUGAUAU | AOAOCHOC CCCHTCTT | |

Where "X" represents stem II region of a HH ribozyme (Hertel et al., 1992 <u>Nucleic Acids Res.</u> 20 3252). The length of stem II may be \geq 2 base-pairs.

Table VI: Potato Citrate Synthase Hairpin Ribozyme and Target Sequences

| Nt. | Ribozyme | The second secon | Substrate |
|-------|--------------------------------------|--|--|
| Posi- | NIDOZ JMC | | and the second of the second o |
| tion | | • | |
| 15 | CAAGUA AGAA GAUG ACCAGAGAAACACACGUUG | UGGUACAUUACCUGGUA | CAUCA GCC UACUUG |
| 112 | CUCGAG AGAA GAGC ACCAGAGAAACACACGUUG | UGGUACAUUACCUGGUA | GCUCC GCU CUCGAG |
| 123 | CUGUUG AGAA GCUC ACCAGAGAAACACACGUUG | UGGUACAUUACCUGGUA | GAGCG GUC CAACAG |
| 181 | GAUCAA AGAA GGAA ACCAGAGAAACACACGUUG | SUGGUACAUUACCUGGUA | UUCCG GUC UUGAUC |
| 285 | CAUAUC AGAA GUGA ACCAGAGAAACACACGUUC | UGGUACAUUACCUGGUA | UCACA GUU GAUAUG |
| 354 | HCCCHC AGAA GGGU ACCAGAGAAACACACGUUG | SUGGUACAUUACCUGGUA | ACCCU GAU GAGGGA |
| 539 | AUGAUG AGAA GGGA ACCAGAGAAACACACGUUG | GUGGUACAUUACCUGGUA | UCCCU GAU CAUCAU |
| 579 | UGGAUG AGAA GUGA ACCAGAGAAACACACGUUG | GUGGUACAUUACCUGGUA | UCACA GCU CAUCCA |
| 596 | GUAGCA AGAA GGGU ACCAGAGAAACACACGUUG | GUGGUACAUUACCUGGUA | ACCCA GUU UGCUAC |
| 710 | UGAGCA AGAA GAUU ACCAGAGAAACACACGUU | | AAUCU GAU UGCUCA |
| 735 | AACAUA AGAA GCAA ACCAGAGAAACACACGUU | | UUGCU GCU UAUGUU |
| 945 | GUANGG AGAA GACA ACCAGAGAAACACACGUUG | GUGGUACAUUACCUGGUA | UGUCU GAU CCUUAC |
| 966 | CAAAGC AGAA GCAA ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | UUGCU GCU GCUUUG |
| 969 | AUUCAA AGAA GCAG ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | CUGCU GCU UUGAAU |
| 988 | GAAGUG AGAA GGCU ACCAGAGAAACACACGUU | | AGCCG GAC CACUUC |
| 1038 | UUCUAC AGAA GAUU ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | AAUCU GUU GUAGAA |
| 1076 | UCUUUC AGAA GCUC ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | GAGCA GUU GAAAGA |
| 1214 | AGUUGA AGAA GUGG ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | CCACU-GUU UCAACU |
| 1253 | IIGUAAG AGAA GGAG ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | CUCCU GUU CUUACA |
| 1356 | AAAGAG AGAA GUAU ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | AUACG GUC CUCUUU |
| 1583 | HIGAAG AGAA GACC ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | GGUCG GCC CUUCAA |
| 1609 | ACTUUG AGAA GAAG ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | CUUCA GUU CAAACU |
| 1720 | UGCUGG AGAA GCGU ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | ACGCU GAU CCAGCA |
| 1819 | UGANIC AGAA GAUU ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | AAUCG GCU GAUUCA |
| 1822 | GCGUGA AGAA GCCG ACCAGAGAAACACACGUU | GUGGUACAUUACCUGGUA | CGGCU GAU UCACCIO |

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Claims

- An enzymatic nucleic acid molecule with RNA cleaving activity, wherein said nucleic acid molecule modulates the expression of a plant gene involved in the biosynthesis of alkaloid compounds.
 - 2. The enzymatic nucleic acid molecule of claim 1, wherein said plant is a solanaceous plant.
- The enzymatic nucleic acid molecule of claim
 wherein said plant is selected from a group consisting of potato, tomato, pepper, eggplant and ditura.
- 4. The enzymatic nucleic acid molecule of claim 15 1, wherein said nucleic acid is in a hammerhead configuration.
- 5. The enzymatic nucleic acid molecule of claim 1, wherein said nucleic acid is in a hairpin 20 configuration.
- 6. The enzymatic nucleic acid molecule of claim 1, wherein said nucleic acid is in a hepatitis δ virus, group I intron, group II intron, VS nucleic acid or 25 RNaseP nucleic acid configuration.
 - 7. The enzymatic nucleic acid of claim 1, wherein said nucleic acid comprises between 12 and 100 bases complementary to RNA of said gene.
 - 8. The enzymatic nucleic acid of claim 1, wherein said nucleic acid comprises between 14 and 24 bases complementary to RNA of said gene.

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- 9. The enzymatic nucleic acid of claim 4, wherein said hammerhead comprises a stem II region of length greater than on equal to two base-pairs.
- 5 10. The enzymatic nucleic acid of claim 5, wherein said hairpin comprises a stem II region of length between three and seven base-pairs.
- 11. The enzymatic nucleic acid of claim 5, wherein said hairpin comprises a stem IV region of length greater than or equal to two base-pairs.
- 12. The enzymatic nucleic acid of claim 1, wherein said gene is solanidine UDP-glucose glucosyl-transferase.
- 13. The enzymatic nucleic acid molecule of claim 12, wherein said nucleic acid specifically cleaves any of sequences shown in Table III, wherein said nucleic acid is in a hammerhead configuration.
 - 14. The enzymatic nucleic acid molecule of claim 12, wherein said nucleic acid specifically cleaves any of sequences shown in Table IV, wherein said nucleic acid is in a hairpin configuration.
 - 15. The enzymatic nucleic acid molecule of any of claims 13 or 14, consisting essentially of one or more sequences selected from the group shown in Tables III and IV.
 - 16. A plant cell comprising the enzymatic nucleic acid molecule of claim 1.

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- 17. A transgenic plant and the progeny thereof, comprising the enzymatic nucleic acid molecule of claim 1.
- 18. An expression vector comprising nucleic acid encoding the enzymatic nucleic acid molecule of claim 1, in a manner which allows expression and/or delivery of that enzymatic nucleic acid molecule within a plant cell.
- 10 19. An expression vector comprising nucleic acid encoding a plurality of enzymatic nucleic acid molecules of claim 1, in a manner which allows expression and/or delivery of said enzymatic nucleic acid molecules within a plant cell.

20. A plant cell comprising the expression vector of claim 18.

- 21. A plant cell comprising the expression vector 20 of claim 19.
 - 22. A transgenic plant and the progeny thereof, comprising the expression vector of claim 18.
- 23. A transgenic plant and the progeny thereof, comprising the expression vector of claim 19.
- 24. A method for modulating expression of an gene in a plant by administering to said plant the enzymatic30 nucleic acid molecule of claim 1.
 - 25. The method of claim 24, wherein said plant is a potato plant.

- 26. The method of claim 24, wherein said gene is solanidine UDP-glucose glucosyl-transferase.
- 27. The expression vector of claim 18, wherein said vector comprises:
 - a) a transcription initiation region;
 - b) a transcription termination region;
 - c) a gene encoding at least one said enzymatic nucleic acid molecule; and
- wherein said gene is operably linked to said initiation region and said termination region, in a manner which allows expression and/or delivery of said enzymatic molecule within said plant cell.
- 15 28. The expression vector of claim 18, wherein said vector comprises:
 - a) a transcription initiation region;
 - b) a transcription termination region;
 - c) an open reading frame;
- d) a gene encoding at least one said enzymatic nucleic acid molecule, wherein said gene is operably linked to the 3'-end of said open reading frame; and

wherein said gene is operably linked to said initiation region, said open reading frame and said termination region, in a manner which allows expression and/or delivery of said enzymatic molecule within said plant cell.

- 29. The expression vector of claim 18, wherein 30 said vector comprises:
 - a) a transcription initiation region;
 - a transcription termination region;
 - c) an intron;

d) a gene encoding at least one said enzymatic nucleic acid molecule; and

wherein said gene is operably linked to said initiation region, said intron and said termination region, in a manner which allows expression and/or delivery of said enzymatic molecule within said plant cell.

- 30. The expression vector of claim 18, wherein 10 said vector comprises:
 - a transcription initiation region;
 - b) a transcription termination region;
 - c) an intron;
 - d) an open reading frame;
- e) a gene encoding at least one said enzymatic nucleic acid molecule, wherein said gene is operably linked to the 3'-end of said open reading frame; and

wherein said gene is operably linked to said initiation region, said intron, said open reading frame 20 and said termination region, in a manner which allows expression and/or delivery of said enzymatic molecule within said plant cell.

- 31. A transgenic plant comprising nucleic acid molecule encoding for an enzymatic nucleic acid molecule with RNA cleaving activity, wherein said nucleic acid molecule modulates the expression of a gene involved in the biosynthesis of alkaloid in said plant.
- 30 32. The transgenic plant of Claim 31, wherein said gene is solanidine UDP-glucose glucosyl-transferase.
 - 33. The transgenic plant of Claim 31, wherein the plant is transformed with Agrobacteriurn, bombarding with

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DNA coated microprojectiles, whiskers, or electroporation.

- 34. The transgenic plant of Claim 33, wherein said bombarding with DNA coated microprojectiles is done with the gene gun.
- 35. The transgenic plant of Claim 31, wherein said plant contains a selectable marker selected from the group consisting of chlorosulfuron, hygromycin, bar gene, bromoxynil, and kanamycin and the like.
- 36. The transgenic plant of Claim 31, wherein said nucleic acid is operably linked to a promoter selected from the group consisting of octopine synthetase, the nopaline synthase, the manopine synthetase, cauliflower mosaic virus (35S); ribulose-1, 6-biphosphate (RUBP) carboxylase small subunit (ssu), the beta-conglycinin, the phaseolin promoter, napin, gamma zein, globulin, the ADH promoter, heat-shock, actin, and ubiquitin.
- 37. The transgenic plant of Claim 31, said enzymatic nucleic acid molecule is in a hammerhead, hairpin, hepatitis Δ virus, group I intron, group II intron, 25 VS nucleic acid or RNaseP nucleic acid configuration
 - 38. The transgenic plant of Claim 31, wherein said enzymatic nucleic acid with RNA cleaving activity encoded as a monomer.

39. The transgenic plant of Claim 31, wherein said enzymatic nucleic acid with RNA cleaving activity encoded as a multimer.

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40. The transgenic plant of Claim 31, wherein the nucleic acids encoding for said enzymatic nucleic acid molecule with RNA cleaving activity is operably linked to the 3' end of an open reading frame.

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41. An enzymatic nucleic acid molecule with RNA cleaving activity, wherein said nucleic acid molecule modulates the expression of a plant gene involved in the flower formation.

- 42. The enzymatic nucleic acid molecule of claim 41, wherein said plant is a potato plant.
- 43. The enzymatic nucleic acid molecule of claim 15 41, wherein said plant is selected from a group consisting of Lettuce, spinach, cabbage, brussel sprouts, arugula, kale, collards, chard, beet, turnip, sweet potato and turfgrass.
- 20 44. The enzymatic nucleic acid molecule of claim 41, wherein said nucleic acid is in a hammerhead configuration.
- 45. The enzymatic nucleic acid molecule of claim 25 41, wherein said nucleic acid is in a hairpin configuration.
- 46. The enzymatic nucleic acid molecule of claim 41, wherein said nucleic acid is in a hepatitis δ virus, 30 group I intron, group II intron, VS nucleic acid or RNaseP nucleic acid configuration.

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- 47. The enzymatic nucleic acid of claim 41, wherein said nucleic acid comprises between 12 and 100 bases complementary to RNA of said gene.
- 5 48. The enzymatic nucleic acid of claim 41, wherein said nucleic acid comprises between 14 and 24 bases complementary to RNA of said gene.
- 49. The enzymatic nucleic acid of claim 44, 10 wherein said hammerhead comprises a stem II region of length greater than on equal to two base-pairs.
- 50. The enzymatic nucleic acid of claim 45, wherein said hairpin comprises a stem II region of length between three and seven base-pairs.
 - 51. The enzymatic nucleic acid of claim 45, wherein said hairpin comprises a stem IV region of length greater than or equal to two base-pairs.
 - 52. The enzymatic nucleic acid of claim 41 wherein said gene is citrate synthase.
- 53. The enzymatic nucleic acid molecule of claim 25 52, wherein said nucleic acid specifically cleaves any of sequences shown in Table V, wherein said nucleic acid is in a hammerhead configuration.
- 54. The enzymatic nucleic acid molecule of claim 30 52, wherein said nucleic acid specifically cleaves any of sequences shown in Table VI, wherein said nucleic acid is in a hairpin configuration.

55. The enzymatic nucleic acid molecule of any of claims 53 or 54, consisting essentially of one or more sequences selected from the group shown in Tables V and VI.

- 56. A plant cell comprising the enzymatic nucleic acid molecule of claim 41.
- 57. A transgenic plant and the progeny thereof, 10 comprising the enzymatic nucleic acid molecule of claim 41.
- 58. An expression vector comprising nucleic acid encoding the enzymatic nucleic acid molecule of claim 41, in a manner which allows expression and/or delivery of that enzymatic nucleic acid molecule within a plant cell.
- 59. An expression vector comprising nucleic acid encoding a plurality of enzymatic nucleic acid molecules of claim 41, in a manner which allows expression and/or delivery of said enzymatic nucleic acid molecules within a plant cell.
- 60. A plant cell comprising the expression vector 25 of claim 58.
 - 61. A plant cell comprising the expression vector of claim 59.
- 30 62. A transgenic plant and the progeny thereof, comprising the expression vector of claim 58.
 - 63. A transgenic plant and the progeny thereof, comprising the expression vector of claim 59.

64. A method for modulating expression of an gene in a plant by administering to said plant the enzymatic nucleic acid molecule of claim 41.

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- 65. The method of claim 64, wherein said plant is a potato plant.
- 66. The method of claim 64, wherein said gene is 10 citrate synthase.
 - 67. The expression vector of claim 58, wherein said vector comprises:
 - a) a transcription initiation region;
- b) a transcription termination region;
 - c) a gene encoding at least one said enzymatic nucleic acid molecule; and

wherein said gene is operably linked to said initiation region and said termination region, in a 20 manner which allows expression and/or delivery of said enzymatic molecule within said plant cell.

- 68. The expression vector of claim 58, wherein said vector comprises:
- 25 a) a transcription initiation region;
 - b) a transcription termination region;
 - c) an open reading frame;
 - d) a gene encoding at least one said enzymatic nucleic acid molecule, wherein said gene is operably linked to the 3'-end of said open reading frame; and
- wherein said gene is operably linked to said initiation region, said open reading frame and said termination region, in a manner which allows expression

and/or delivery of said enzymatic molecule within said plant cell.

- 69. The expression vector of claim 58, wherein 5 said vector comprises:
 - a transcription initiation region;
 - b) a transcription termination region;
 - c) an intron;
- d) a gene encoding at least one said enzymatic
 10 nucleic acid molecule; and

wherein said gene is operably linked to said initiation region, said intron and said termination region, in a manner which allows expression and/or delivery of said enzymatic molecule within said plant cell.

- 70. The expression vector of claim 58, wherein said vector comprises:
 - a) a transcription initiation region;
 - b) a transcription termination region;
 - c) an intron;
 - d) an open reading frame;
- e) a gene encoding at least one said enzymatic nucleic acid molecule, wherein said gene is operably linked to the 3'-end of said open reading frame; and
 - wherein said gene is operably linked to said initiation region, said intron, said open reading frame and said termination region, in a manner which allows expression and/or delivery of said enzymatic molecule
- 30 within said plant cell.
 - 71. A transgenic plant comprising nucleic acid molecule encoding for an enzymatic nucleic acid molecule with RNA cleaving activity, wherein said nucleic acid

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molecule modulates the expression of a gene involved in flower formation in said plant.

- 72. The transgenic plant of Claim 71, wherein said 5 gene is citrate synthase.
- 73. The transgenic plant of Claim 71, wherein the plant is transformed with Agrobacteriurn, bombarding with DNA coated microprojectiles, whiskers, or electroporation.
 - 74. The transgenic plant of Claim 73, wherein said bombarding with DNA coated microprojectiles is done with the gene gun.

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75. The transgenic plant of Claim 71, wherein said plant contains a selectable marker selected from the group consisting of chlorosulfuron, hygromycin, bar gene, bromoxynil, and kanamycin and the like.

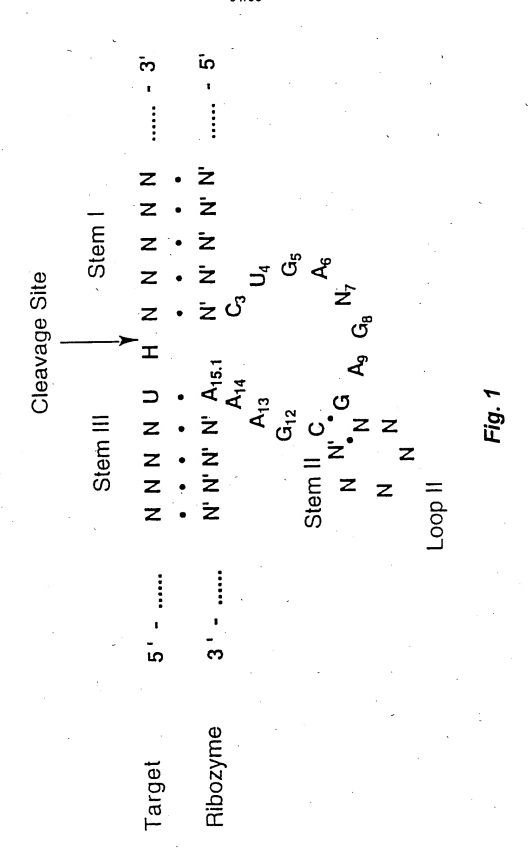
- 76. The transgenic plant of Claim 71, wherein said nucleic acid is operably linked to a promoter selected from the group consisting of octopine synthetase, the nopaline synthase, the manopine synthetase, cauliflower mosaic virus (35S); ribulose-1, 6-biphosphate (RUBP) carboxylase small subunit (ssu), the beta-conglycinin, the phaseolin promoter, napin, gamma zein, globulin, the ADH promoter, heat-shock, actin, and ubiquitin.
- 30 77. The transgenic plant of Claim 71, said enzymatic nucleic acid molecule is in a hammerhead, hairpin, hepatitis δ virus, group I intron, group II intron, VS nucleic acid or RNaseP nucleic acid configuration

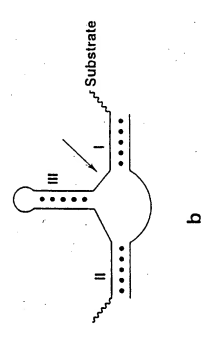
The transgenic plant of Claim 71, wherein said enzymatic nucleic acid with RNA cleaving activity encoded as a monomer.

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The transgenic plant of Claim 71, wherein said enzymatic nucleic acid with RNA cleaving activity encoded as a multimer.

The transgenic plant of Claim 71, wherein the 10 nucleic acids encoding for said enzymatic nucleic acid molecule with RNA cleaving activity is operably linked to the 3' end of an open reading frame.





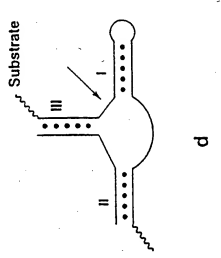
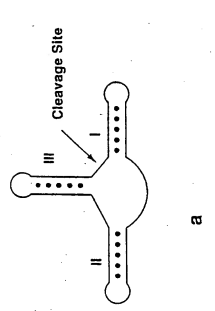
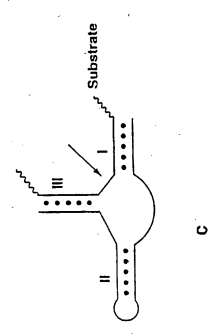
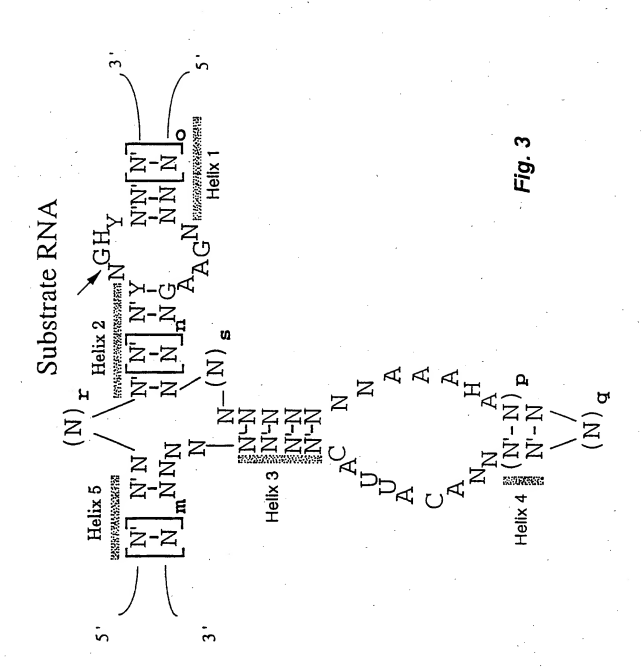


Fig. 2

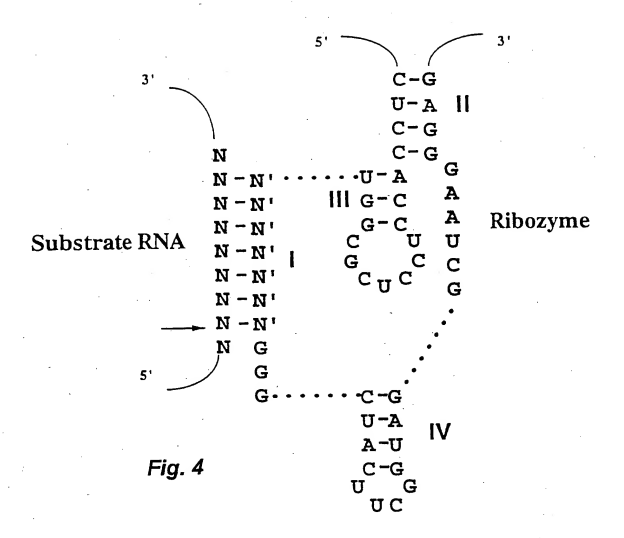




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